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# 1 Reversible superdense ordering of lithium between two graphene 2 sheets

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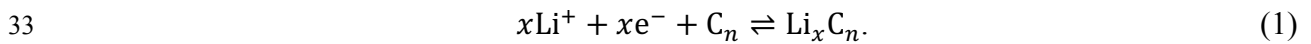
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13 **Many carbon allotropes can act as host materials for reversible lithium uptake,<sup>1,2</sup> laying**  
14 **the grounds for existing and future electrochemical energy storage. Insights into exactly**  
15 **how lithium is arranged within those are, however, challenging to obtain *in operando*.**  
16 **Using *in situ* transmission electron microscopy (TEM)<sup>3-5</sup> to probe light elements (especially**  
17 **Li)<sup>6,7</sup> is severely hampered by their low scattering cross section for impinging electrons**  
18 **and their susceptibility to knock-on damage<sup>8</sup>. Here, we study the reversible Li intercalation**  
19 **of bilayer graphene by *in situ* spherical and chromatic aberration-corrected low-voltage**  
20 **TEM<sup>9</sup>, supported by electron energy loss spectroscopy (EELS) and density functional**  
21 **theory (DFT) calculations. Upon their remote insertion from an electrochemical gate, we**  
22 **observe Li atoms to assume multi-layered close-packed order between the two carbon**

23 **sheets. The associated Li storage capacity by far exceeds the one of LiC<sub>6</sub>—the densest**  
24 **configuration known under normal conditions for bulk graphitic carbon<sup>10</sup>. Our findings**  
25 **thus point to the possible existence of distinct storage arrangements of ions in 2D layered**  
26 **materials as compared to their bulk parent compounds.**

27 Figure 1a shows a schematic of our devices, all of which are supported by Si<sub>3</sub>N<sub>4</sub>-covered Si  
28 substrates. The Si<sub>3</sub>N<sub>4</sub> forms a 40×40 μm<sup>2</sup> membrane at the centre of the Si-chip. The bilayer  
29 graphene flake is exfoliated from natural graphite and etched into a Hall bar shape. One end of  
30 the flake is connected to a counter electrode *via* a Li-ion conducting solid polymer electrolyte.  
31 This setup allows for the controlled electrochemical reduction/oxidation of bilayer graphene  
32 according to<sup>1</sup>



34 Similar to Refs. [11, 12], we trigger lithiation (delithiation) by applying a positive voltage  $U_G =$   
35  $5 \text{ V}$  ( $U_G = 0 \text{ V}$ ) to the counter electrode with respect to bilayer graphene. A grounded current  
36 lead to the latter serves as a source/sink for electrons required to facilitate the reversible  
37 intercalation of Li-ions at the electrolyte-covered end of bilayer graphene. Intercalated Li  
38 exhibits rapid lateral diffusion that tends to establish and maintain an even distribution of Li  
39 throughout the bilayer.<sup>13</sup> Hence, one may study its ordering in a region well separated from the  
40 electrolyte by *in situ* TEM (Figs. 1a–b), thereby also preventing exposure of the electrolyte to  
41 the electron beam. In the region probed by TEM, bilayer graphene is suspended over a hole in  
42 the Si<sub>3</sub>N<sub>4</sub> membrane. Metallic contacts to the bilayer allow monitoring its resistivity  $\rho_{xx} =$   
43  $U_{xx}/I$  in the electrolyte-uncovered region (Fig. 1a). During subsequent lithiation/delithiation  
44 cycles, we typically observe reversible changes in  $\rho_{xx}$  (Fig. 1c). These relate to changes in the  
45 local Li concentration *via* finite electronic charge transfer.<sup>12,13</sup> The drop in  $\rho_{xx}$  during lithiation  
46 reflects an increase in electron density, characteristic for ambipolar diffusion of electron-ion

47 pairs into the probed area.<sup>13</sup> During delithiation, Li-ions and electrons leave the bilayer, thereby  
48 restoring its initial resistivity value. The exact time evolution of  $\rho_{xx}$  depends on the kinetics of  
49 several (uncontrolled) processes, related to e.g. ionic transport within the electrolyte and across  
50 the solid electrolyte interphase. Yet, Fig. 1c is a qualitative characteristic of reversible Li  
51 intercalation in bilayer graphene.<sup>13</sup> In the following, we present *in situ* TEM data obtained in the  
52 *SALVE* (Sub-Ångström Low-Voltage Electron microscopy) microscope<sup>9</sup>. We work at an  
53 electron acceleration voltage of 80 kV, just below the threshold for knock-on damage of C atoms  
54 in graphene.<sup>14</sup> Despite these conditions, the instrument still delivers sub-Ångström resolution in  
55 the images. Further details can be found in Methods (also Extended Data Figs. 1–2).

56  
57 Figure 2a shows a TEM image of pristine bilayer graphene. The inset depicts its Fourier  
58 transform (FT) and confirms the known value of the in-plane lattice constant  $a_C = 2.46 \text{ \AA}$ . The  
59 image marks the beginning of a series acquired during lithiation of a bilayer graphene device  
60 (Figs. 2a–c and Supplementary Video 1), but it is representative of the sample state prior to  
61 application of the bias voltage  $U_G = 5 \text{ V}$ . In Fig. 2b, acquired after 170 s, a second crystal lattice  
62 has appeared in the lower half of the probed area. White dashed lines demarcate its boundary on  
63 the left and right of the image. The image in Fig. 2c is recorded at  $t = 288 \text{ s}$ . The additional  
64 crystal structure now extends throughout almost the whole field of view. Fig. 2d is the Fourier  
65 transform of Fig. 2b. Compared with the Fourier transform of pristine bilayer graphene (inset to  
66 Fig. 2a), we identify three sets of additional signals, highlighted in red, green, and blue. These  
67 attest to hexagonal crystalline order (as for graphene), but with an in-plane lattice constant of  
68  $3.1 \text{ \AA}$ . In Fig. 2e, their spatial distribution is mapped. This allows discerning three grains none  
69 of which are aligned to the encapsulating graphene lattice. In Fig. 2f the same Fourier transform  
70 is shown but with a von Hann filter applied to minimize the streaks. The highlighted signals  
71 stem from both bilayer graphene (cyan) and the additional crystalline phase (green) as well as

72 Moiré artefacts (magenta) and their origin (bold arrows). Fig. 2g is a Fourier-filtered (Methods  
73 and Extended Data Fig. 3) version of panel b, where the graphene lattice, as well as the Moiré  
74 effects, have been removed (see panels h and i for a magnification before and after filtering).  
75 These images offer a direct view on the encapsulated crystal. The observed contrast in the images  
76 also suggests regions of different thickness even within a single grain. This is worked out in  
77 detail in Methods and Extended Data Fig. 4.

78

79 To narrow down the chemical composition of the additional crystal structure, EELS data has  
80 been acquired on bilayer graphene before and during lithiation (Fig. 2j). Before lithiation, we  
81 exclusively observe the C K-edge at 284 eV. During lithiation the Li K-edge at 55 eV on regions  
82 characterized by a Fourier transform as in Fig. 2d is additionally detected. Based on the absence  
83 of other signals in the explored energy range of 0–800 eV, Si (L<sub>2,3</sub>-edge starting at 99 eV), S  
84 (L<sub>2,3</sub>-edge starting at 165 eV), N (K-edge starting at 400 eV), O (K-edge starting at 532 eV and  
85 F (K-edge starting at 685 eV) are discarded as significant constituents of the new crystalline  
86 phase. Likewise, Ti and Pt (electrode material) can be ruled out based on the observed light  
87 atomic contrast in Fig. 2 for the case of Pt and on the absence of the distinct Ti L<sub>2,3</sub>-edge at 456  
88 eV in the EELS data. Although it is inherently impossible to exclude H or C, the crystalline  
89 phase formed during lithiation likely consists of pure Li. The low onset in energy of the Li K-  
90 edge supports this assertion.<sup>15,16</sup> Also, a Li plasmon mode appears near 9 eV<sup>17,18</sup> (Extended Data  
91 Fig. 5a). Though the shape of the Li-K edge resembles the one of Li<sub>2</sub>O/LiOH,<sup>15,16</sup> stoichiometric  
92 Li<sub>2</sub>O/LiOH can be disqualified, since we would then expect both a more pronounced O-K edge  
93 as well as an imaging contrast comparable in strength to the encapsulating graphene.<sup>19</sup> Good  
94 agreement with experiment is attained when calculating the Li-K edge shape of graphene-  
95 encapsulated Li multilayers (Methods and Extended Data Fig. 5b). We do not rule out the  
96 presence of trace oxygen, also suggested by the occasional observation of a weak shoulder near

97 30 eV (Extended Data Fig. 5a), previously attributed to oxidized lithium.<sup>17,18</sup> Yet, the extracted  
98 in-plane lattice constant of 3.1 Å matches close-packed Li.<sup>20</sup> This coincidence is surprising,  
99 since normally very low temperatures and/or extreme pressures are required for Li to assume  
100 this ordering.<sup>21</sup>

101

102 To test whether indeed the formation of a dense, multi-layered Li crystal in bilayer graphene is  
103 conceivable, first-principles calculations were carried out (Methods). The chemical potential of  
104 Li atoms in bulk close-packed phases was evaluated. All energies below are given with respect  
105 to the hexagonal close-packed (hcp) phase, which had the lowest energy in the calculations.  
106 Several layers of Li atoms inside bilayer graphene are considered (Fig. 3 and Extended Data  
107 Figs. 6–8). Since the *in situ* TEM studies do not reveal evidence for a change in registry of the  
108 graphene sheets (Extended Data Fig. 9), AB stacking is assumed as in the pristine device. Our  
109 main conclusions, however, hold irrespective of the stacking order. The relaxation of atomic  
110 coordinates of a single layer of Li atoms yields an energetically favoured C<sub>6</sub>LiC<sub>6</sub> configuration,  
111 with Li arranged in a commensurate ( $\sqrt{3}\times\sqrt{3}$ )R30° superstructure with a lattice constant of  $a_{\text{LiC}_6}$   
112 = 4.26 Å (Figs. 3a–b). Except for the graphene registry, this finding is similar to the bulk LiC<sub>6</sub>  
113 phase<sup>10</sup> and the AA-stacked C<sub>6</sub>LiC<sub>6</sub> phase<sup>22</sup>. The situation changes for a larger number of Li  
114 layers. Finite clusters and infinite (periodic) hcp structures with different orientation with respect  
115 to the graphene lattice were considered (Figs. 3c–f, Extended Data Figs. 7–8). The energies of  
116 these systems are very close to that of the C<sub>6</sub>LiC<sub>6</sub> configuration (higher by only 0.01–0.05 eV/  
117 Li atom). The in-plane lattice constant  $a_{\text{Li}}$  of the Li hcp bilayer and trilayer (average distances,  
118 as the positions of Li atoms are affected by nearby C atoms) is in the range of 3.05–3.15 Å,  
119 matching the experimental value of 3.1 Å as well as the identical literature value.<sup>20</sup> These results  
120 suggest that the formation of a multilayer close-packed Li phase between graphene sheets is  
121 conceivable. Its precise stacking order may however differ from hcp as other configurations are

122 energetically similar (Extended Data Fig. 8d–e). These are hard to distinguish in these TEM  
123 experiments, because we may only observe the projection along one crystal direction. Moreover,  
124 some bulk diffraction selection rules do not hold in the given case of an atomically thin  
125 specimen. TEM image simulations reveal comparable contrast and diffraction patterns for the  
126 two extreme cases: cubic close-packed (ccp) and hexagonal close packed (Extended Data Fig.  
127 2f–g).

128

129 By analysing the electronic structure and charge distribution, the charge transfer between Li and  
130 graphene can be estimated. Fig. 3b,d and f display colour renditions of the charge probability  
131 distribution as compared to isolated graphene and Li crystals consisting of a single, double or  
132 triple layer. As can be seen from Fig. 3f in the triple layer case, charge transfer is significant  
133 only for the outer Li layers directly neighbouring a graphene sheet. Inner Li layers retain their  
134 metallic character as electronic charge is distributed between the  $\text{Li}^+$  lattice sites. Extended Data  
135 Fig. 8c depicts the average charge transfer per Li atom, which drops as the close-packed Li phase  
136 gets thicker. Renormalizing by the number of atoms in the outermost Li layers only, we find a  
137 constant charge transfer of approximately  $0.33 \text{ e}^-/\text{outer Li atom}$ , irrespective of how many  
138 (metallic) Li layers (with nearly zero charge transfer) are packed in between. This is consistent  
139 with the observation that the resistivity  $\rho_{xx}$  measured during lithiation tends to saturate rather  
140 than to decrease progressively as more Li enters bilayer graphene (Fig. 1c). Note that although  
141 for  $\text{C}_6\text{LiC}_6$  (Figs. 3a–b) a higher value of  $0.85 \text{ e}^-/\text{Li atom}$  for the charge transfer applies resulting  
142 in an estimated density of transferred electrons of  $2.7 \cdot 10^{14} \text{ cm}^{-2}$  per graphene sheet, the much  
143 denser arrangement of Li atoms in the close-packed structure yields a higher electron density  
144 per graphene sheet of  $4 \cdot 10^{14} \text{ cm}^{-2}$  despite the smaller charge transfer value.

145

146 During lithiation, the close-packed Li phase grows laterally between the graphene sheets (Fig.  
147 4a and Supplementary Video 1). In Fig. 4a, The graphene signals have been Fourier filtered in  
148 order to reveal the Li crystal structure. Three grains of varying in-plane orientation have been  
149 coloured for the sake of illustration. While the grain boundary between the central grain (green)  
150 and the lower right grain (blue) is rather sharp and stable, the one with the left grain (red) appears  
151 more fuzzy and mobile as their orientation nearly matches. Regions of different thickness can  
152 be identified even within a single grain (Extended Data Fig. 4). The specimen is too thin to  
153 reliably extract its exact thickness  $t$  from EELS (we typically obtain  $t/\lambda < 0.1$ , where  $\lambda$  is the  
154 inelastic mean free path of our 80 keV electrons). One may nonetheless determine relative  
155 variations (see Methods). When imaging an extremely thin slab of weakly scattering elements  
156 in a microscope with sufficient resolving power, the imaging contrast does increase with  
157 increasing specimen thickness. This is quantified in the Extended Data Figs. 4e–f, revealing that  
158 thinner parts of the close-packed Li phase tend to be located closer to its perimeter. At the leading  
159 edge, single atoms can be identified (Methods, Extended Data Fig. 1). From the image time  
160 series, a lateral growth rate of the order of 1 Å/s can be extracted. During delithiation the close-  
161 packed Li phase disassembles and gradually disappears (Fig. 4b). Eventually, the pure bilayer  
162 graphene lattice remains behind. The degree of reversibility of this process is limited by the  
163 number of defects in the graphene lattice and their irreversible formation during prolonged  
164 imaging (electron irradiation). Given the combined slow image acquisition (on the order of 1 s)  
165 and low sensitivity to light atoms, Li diffusing at speed in less ordered configuration remains  
166 concealed to the TEM observer. Such processes are nonetheless present in parallel with the close  
167 packing as well as beyond the boundaries of this phase and are likely responsible for the initial  
168 abrupt change in resistivity  $\rho_{xx}$  during lithiation (Fig. 1c).

169



170 We note that the observed crystalline phase of Li proves stable between intact graphene planes  
171 only. With the incident electron energy exceeding the threshold for displacement damage of Li  
172 ( $\sim 20$  keV), conditions are such that Li readily “boils” under the electron beam<sup>16,18</sup>. The close-  
173 packed phase is volatile when imaged near bilayer edges or in the presence of a high density of  
174 defects in the graphene lattice. Constituents may escape from between graphene sheets *via* such  
175 edges or defects upon electron-beam induced melting of the crystalline phase. Likewise, we find  
176 material having escaped from within and agglomerated on the outer bilayer surfaces near such  
177 defects to quickly evaporate under electron beam irradiation. The protective encapsulation by  
178 two impermeable atomic sheets may thus be regarded a prerequisite to safe probing by TEM of  
179 the crystal formation therein, akin to the situation in graphene liquid cells<sup>23</sup>.

180

181 Close-packing of Li intercalated between graphene sheets, as demonstrated here, results in a  
182 structure with Li-content in great excess of  $\text{LiC}_6$ . Although enhanced Li storage has been  
183 proposed to occur on the outside of graphene planes<sup>24,25</sup> previously, suggested atomistic  
184 configurations were contradictory among the different works and could neither be addressed nor  
185 verified by microscopic means. The same holds for the scarce reports claiming the formation of  
186 nanocrystallites of close-packed Li during the lithiation of different carbon allotropes.<sup>26,27</sup> At  
187 elevated temperatures, a similar configuration of Li, as we report here, may, as a matter of fact,  
188 have been left unidentified in bulk graphite.<sup>28</sup> Since the energy cost for close-packing of Li in  
189 the van der Waals gap of bilayer graphene as compared to forming  $\text{C}_6\text{LiC}_6$  is very similar (Fig.  
190 3), close-packing may be the answer of the system for having to accommodate a larger amount  
191 of Li supplied within a short time. The activation energy for Li diffusion in bilayer graphene has  
192 been calculated (Methods). It points toward facile diffusion of Li between graphene sheets, even  
193 in the presence of a close-packed Li phase. Even though, a bilayer can be regarded as the  
194 fundamental unit of bulk graphite, it may still feature considerably distinct properties and in

195 particular its layers may spread more easily to accommodate for dense intercalate ordering. It  
196 should also be noted that Li-graphite intercalation compounds only occupy a small region in the  
197 C-Li binary alloy phase diagram.<sup>29</sup> In the miscibility gap beyond LiC<sub>6</sub>, alternative configurations  
198 may be available for storing larger amounts of lithium in layered carbons.<sup>30</sup>

199

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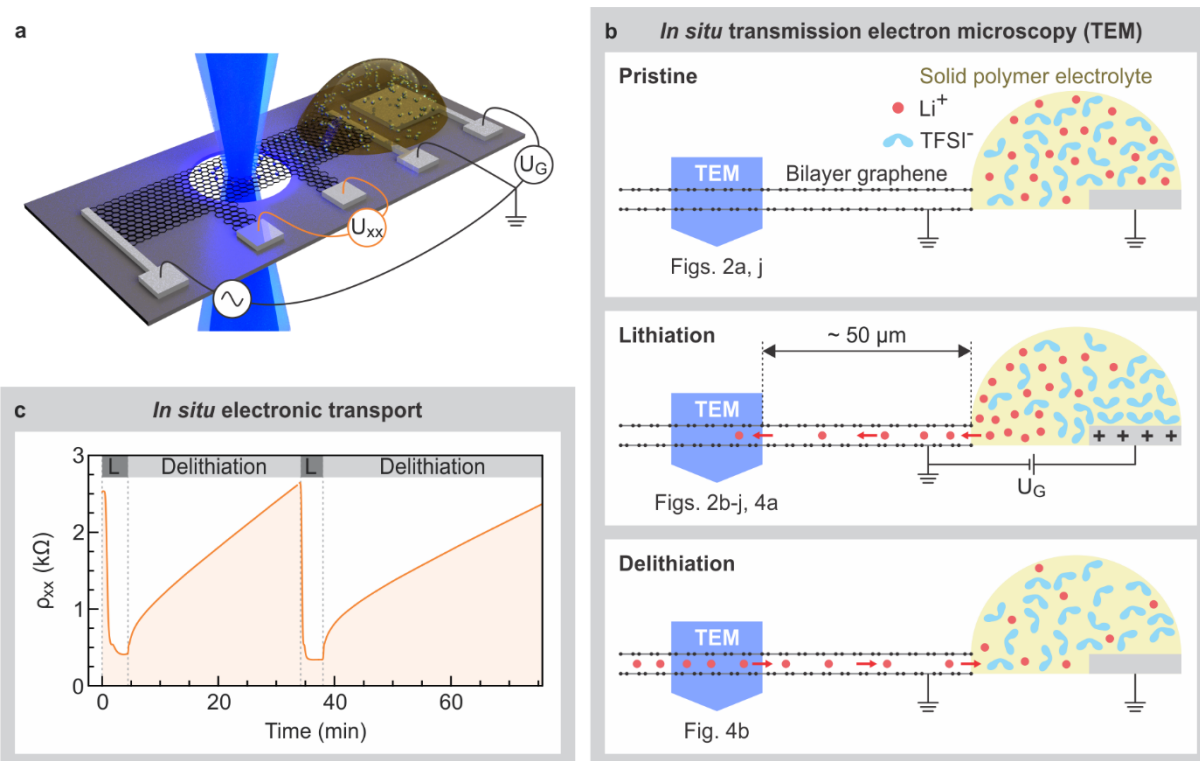
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264

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276 **Author Contributions** J. H. S and U. K. composed the project. M. K. and S. F. fabricated the  
277 samples and performed the electrochemical measurements. F. B. performed the TEM and EELS  
278 experiments. M. G. and A. V. K. did the DFT calculations. J. B. helped with TEM and EELS  
279 experiments and did TEM imaging simulations. U. K. supervised the TEM work. D. S.  
280 contributed to the electrochemical design of the experiment. M. K. and J. H. S. wrote the  
281 manuscript and all authors contributed to it.

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286

287 **Figure 1 | Device layout and working principle. a**, Schematic of the device (not to scale).

288 Bilayer graphene (black) on a substrate (dark grey) is contacted by several metallic electrodes

289 (bright grey). A Li-ion (white spheres) conducting electrolyte (yellow) connects the bilayer with

290 a metallic counter electrode. On the order of  $50 \mu\text{m}$  away from its electrolyte-covered end,

291 bilayer graphene is partially suspended over a hole allowing for transmission electron

292 microscopy (TEM) investigations (electron beam illustrated in blue). **b**, Schematic side view of

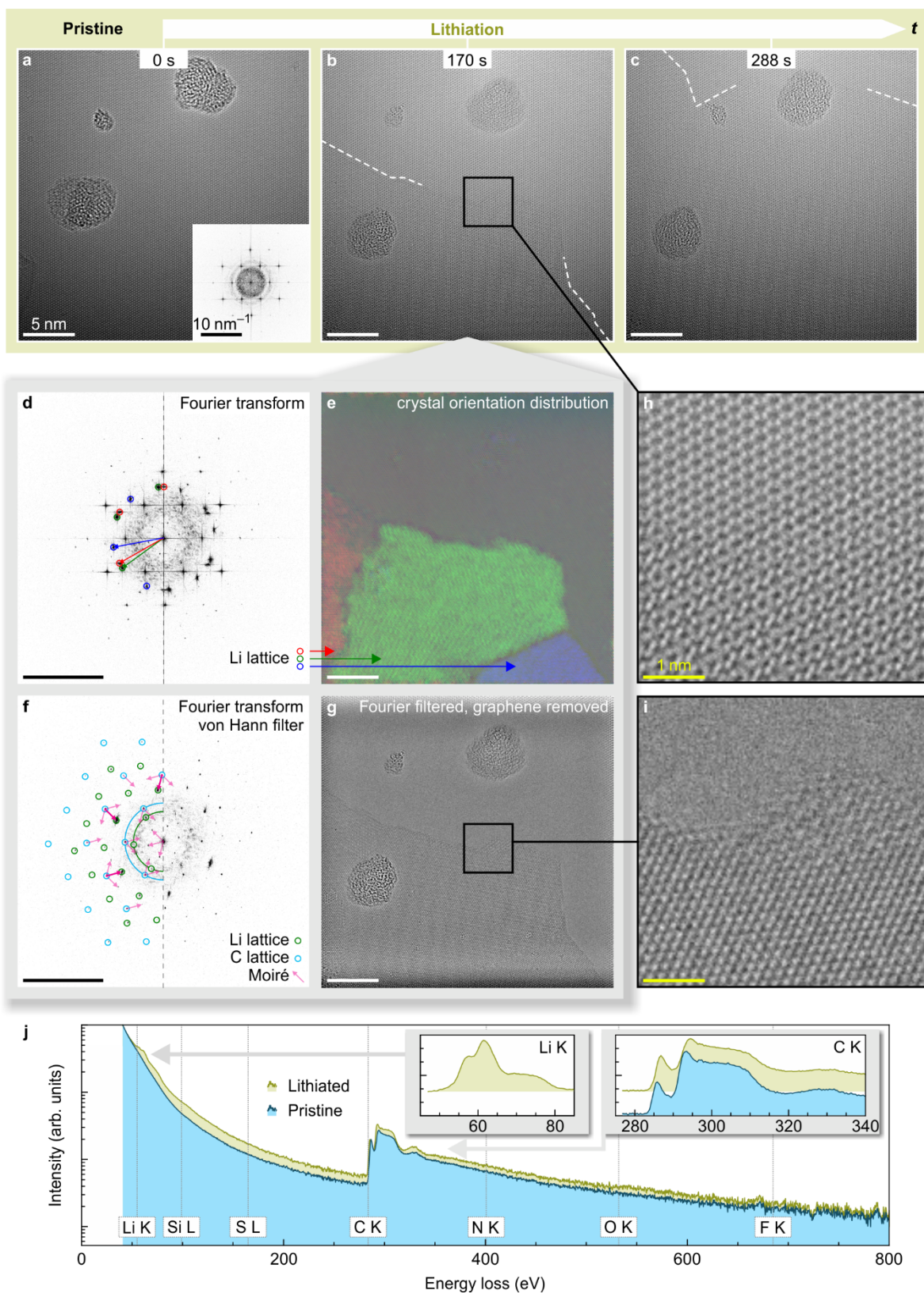
293 the pristine device as well as during lithiation and delithiation. Reference to TEM data acquired

294 at the respective state is given. **c**, Bilayer graphene's resistivity  $\rho_{xx}$  measured *in situ* during two

295 lithiation (L)/delithiation cycles inside the SALVE microscope with the electron beam blanked.

296 As schematically shown in **(a)**, a four point probe configuration serves to reveal reversible

297 changes in the electrolyte-uncovered region of the device.

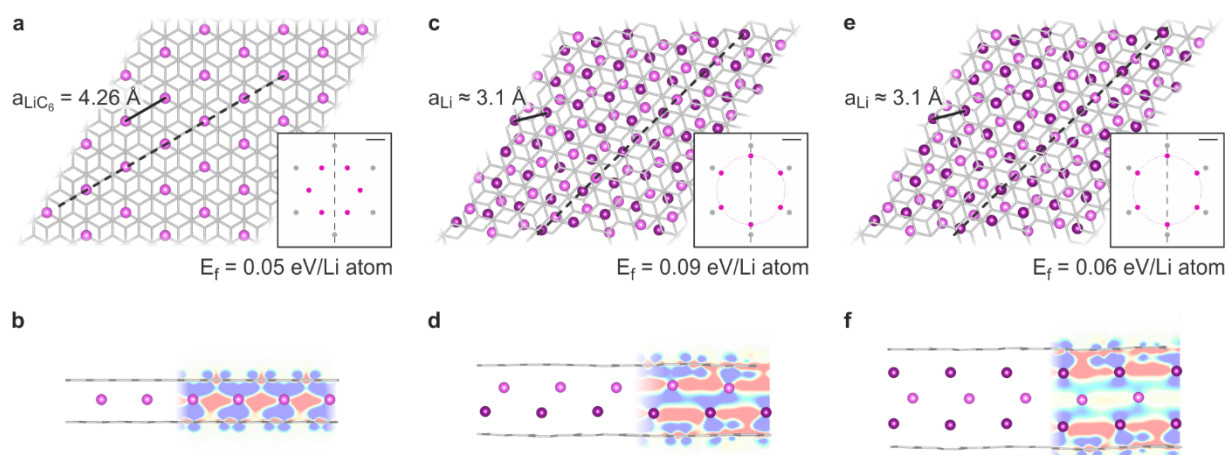


298

299 **Figure 2 | *In situ* TEM measurements.** a–c, TEM images showing the propagating front of a  
 300 Li crystal forming inside bilayer graphene during lithiation. The images are acquired on the



301 same sample area at consecutive times as indicated. Amorphous hydrocarbon adsorbates appear  
302 as few nm-large blobs, located above or below the bilayer. The inset in **(a)** is its Fourier  
303 transform. **d**, Fourier transform of **(b)**. Three sets of spots marked in red, green, and blue are  
304 rotated relative to each other as they stem from three different Li crystal grains. The cross-shaped  
305 streaks are edge artefacts from the Fourier transform. **e**, Spatial distribution of the Li grains in  
306 **(b)** with the colour coding from **(d)**. **f**, von Hann-filtered Fourier transform of **(b)**. Fourier  
307 transforms are point symmetric, therefore, marks on the left side do not mask information.  
308 Signals from bilayer graphene (Li) are highlighted in cyan (green); (the origin of) Moiré artefacts  
309 are highlighted in (bold) magenta. The anisotropic smearing of the Li signals is due to the sharp  
310 propagation front. Signals originating from the grains coloured red and blue in **e** are dampened  
311 away by the applied von Hann filter. Half circles represent the fundamental periodicities of 0.213  
312 nm for graphene (cyan) and 0.276 nm for Li (green). **g**, Fourier-filtered version of **(b)** where the  
313 graphene lattice, as well as the Moiré effects, are filtered out and only the Li crystal structure is  
314 left. The contrast at the edge of the figure is an artefact from the Fourier filter. **h**, **i**, Magnified  
315 detail from the demarcated areas in **b** and **g** respectively, showing the Li crystal edge. Scale bars  
316 of equal size are coloured identically but labelled only once in **(a–g)**. **j**, EELS data with  
317 logarithmic intensity scale before (blue) and during (yellow) lithiation acquired on an area as in  
318 **(a)** and **(c)**, respectively. Highlighted are the energies of the relevant major edges. Insets show  
319 the near-edge structure of the Li and C K-edges on a linear intensity scale and after individual  
320 subtraction of an inverse-power law background. Note that EELS data get noisy at high energies  
321 because of the exponential decay of the signal.



322

323 **Figure 3 | Atomistic models of Li crystals between AB-stacked graphene sheets obtained**

324 **from DFT calculations. a, b, The ‘conventional’  $C_6LiC_6$  configuration with Li arranged in a**

325 **commensurate  $(\sqrt{3}\times\sqrt{3})R30^\circ$  superstructure between graphene sheets. c, d, Fully optimized**

326 **bilayer Li crystal. e, f, Fully optimized trilayer Li crystal (one of two energetically close stacking**

327 **configurations, compare with Extended Data Fig. 8d). The projection of the latter two structures**

328 **matches well the experimental observations. (a, c, e) are top views and (b, d, f) are side views**

329 **along the dashed line given in the respective top view. C atoms and  $sp^2$ -bonds are grey, Li atoms**

330 **are magenta. The insets in (a, c, e) schematically show diffraction patterns associated with the**

331 **respective structure (scale bar is  $2\text{ nm}^{-1}$ ), with first order diffraction spots from the Li (C) lattice**

332 **indicated in magenta (grey). The close-packed Li phase in (c, e) may assume any relative rotation**

333 **angle with respect to the bilayer graphene lattice at virtually no additional energy cost (see**

334 **Extended Data Fig. 7b). A faint magenta circle indicates that the Li diffraction spots may**

335 **therefore be rotated with respect to those of graphene.  $E_f$  is the energy required to take a Li atom**

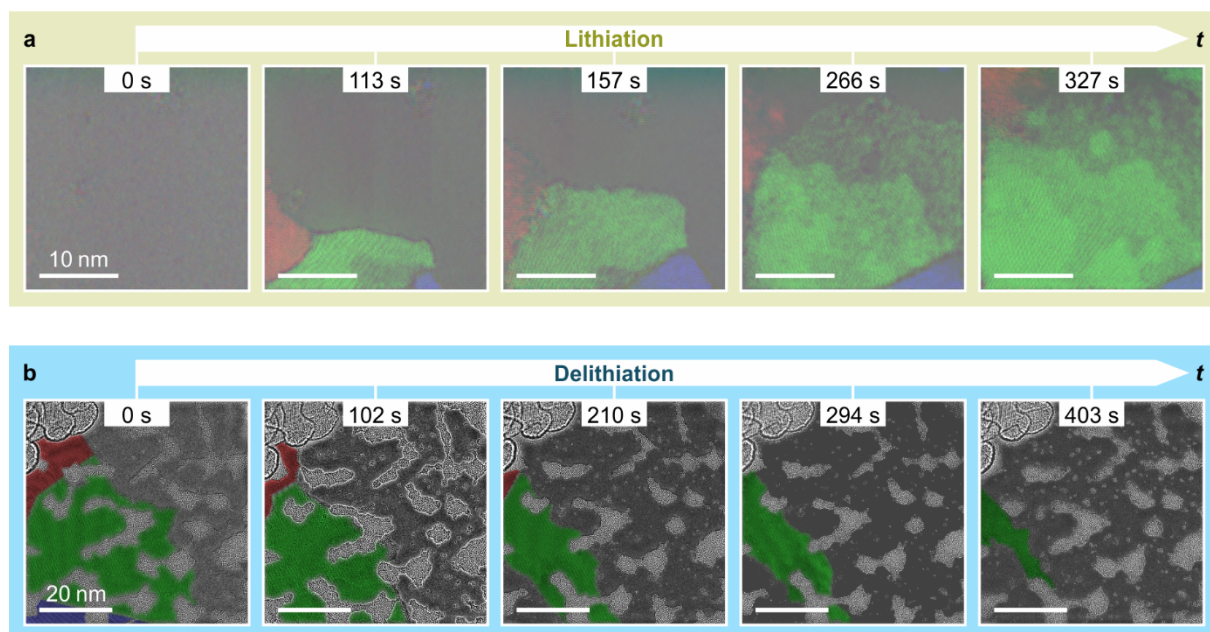
336 **from a bulk Li crystal and insert it between graphene sheets in the corresponding configuration.**

337 **Contour plots in (b, d, f) represent the charge transfer between Li and graphene as compared to**

338 **the isolated graphene and Li crystals. An increase in the electron density (negative charge) is**

339 **blue, and a decrease in the electron density (positive charge) is red. Note the smaller average**

340 **charge transfer to graphene for the multilayer Li crystals.**



341

342 **Figure 4 | Li crystal growth between two graphene sheets. a, b,** Time series of TEM images  
 343 after Fourier filtering of the bilayer graphene signal acquired during (a) lithiation and (b)  
 344 delithiation. Crystal grains of different in-plane orientation are coloured in red, green, and blue.  
 345 An increased amount of amorphous, immobilized residues (grey areas, essentially composed of  
 346 carbon) are present within the field of view in (b), an unavoidable consequence of extended  
 347 electron beam illumination. In both (a, b) the interface with the electrolyte (not contained in the  
 348 images) is oriented towards the bottom of the page.