Flexible and robust MoS$_2$/graphene hybrid paper cross-linked by a polymer ligand: a high-performance anode material for thin film lithium-ion batteries

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S1. Experimental details

S1.1 Raw materials

MoS$_2$ was purchased from Sigma-Aldrich. GO was purchased from Nanjing XF Nano. PEO ($M_w = 100,000$) was purchased from Alfa Aesar. CuCl$_2$·2H$_2$O and DMF were purchased from Beijing Chemical Works. Hydrazine hydrate was purchased from Guangdong Chemical Reagent Engineering-Technological Research and Development Centre.

S1.2 Preparation of a MoS$_2$/DMF solution

The procedures for exfoliating MoS$_2$ into nanosheets were adapted from a previous report [1]. Briefly, the MoS$_2$ powder was added to DMF at an initial concentration of 10 mg mL$^{-1}$, homogenized at 11,500 rpm for 60 min, and then subjected to cup-horn sonication at 300 W for 60 min. The obtained slurry was centrifuged at 1,500 rpm for 45 min, and the top 1/2 supernatant was collected. The collected supernatant was further centrifuged at 3,000 rpm for another 45 min. The precipitated solid was collected and re-dispersed in DMF by sonication, yielding a dark green MoS$_2$/DMF solution (0.2 mg mL$^{-1}$) containing relatively large MoS$_2$ nanosheets (Fig. 1).

S1.3 Preparation of a GO/DMF solution

To produce a GO/DMF solution [2], GO was directly added to DMF at an initial concentration of 0.2 mg mL$^{-1}$, and then subjected to gentle sonication at 100 W for 60 min. The obtained GO/DMF solution was brilliant yellow, and could stand for weeks without obvious precipitates (Fig. S2).

S1.4 Preparation of freestanding MoS$_2$/r-GO paper cross-linked by PEO

The MoS$_2$ and GO/DMF solutions were mixed, and PEO was added at different MoS$_2$/GO/PEO wt ratios
(9/0/1, 7/2/1, 6/3/1 and 5/4/1). A small amount of dehydrated CuCl₂ was added to the MoS₂/GO/PEO system, which was then kept under magnetic stirring for 24 h. After coordination hydrazine hydrate was added, and the MoS₂/GO/PEO system was kept at 80 °C for 12 h such that GO was sufficiently reduced to r-GO. It was reported that GO could be chemically reduced to r-GO by hydrazine hydrate, which would re-stack and thus precipitate from DMF [3]. However, the MoS₂/GO/PEO system could be successfully converted to the MoS₂/r-GO/PEO system by hydrazine hydrate without obvious precipitates (Fig. S7), thanks to the steric stabilisation effect of PEO which facilitated the following liquid-phase processing. The vacuum filtration was employed to obtain a piece of flexible and robust MoS₂/r-GO paper chemically cross-linked by PEO, which was peeled off the poly(vinylidene fluoride) (PVDF) membrane by soaking in acetone (Fig. S8). Note that if the PEO fraction was too low, it was helpless to create flexible and robust MoS₂/r-GO/PEO paper; if the PEO fraction was too high, it would impose an adverse effect on the electrochemical performance of the MoS₂/r-GO/PEO paper due to the decreased electronic conductivity resulting in increased internal resistance and charge transfer [4]. Moreover, it should be mentioned that the higher the r-GO content, the stronger the MoS₂/r-GO/PEO paper. The MoS₂/r-GO/PEO (9/0/1) paper was the most fragile, which would at least partially lead to the poor cycle stability.

**S1.5 Preparation of a film anode for the electrochemical testing**

The freestanding MoS₂/r-GO paper could be tailored arbitrarily by scissors without cracking, and directly attached to a copper foil with a layer of silver paste (Fig. S9). The layer of silver paste was aimed to improve the surface roughness of the copper foil, and thus to increase the efficiency of physical contact between the film anode and the current collector (copper foil) [5]. The film anode was then dried in nitrogen at 120 °C for 12 h to adequately evaporate the residual organic solvent, and equipped in a CR2032 coin cell according to the configuration of (-) Li | electrolyte | freestanding MoS₂/r-GO/PEO film anode (+) with a liquid electrolyte (1 M solution of LiPF₆ in ethylene carbonate–dimethyl carbonate at a vol ratio of 1 : 1) in an
argon-filled glove box. A Celgard 2400 membrane was used as the separator.

**S1.6 Equipment**

TEM was performed by a Hitachi H7700 operated at an accelerating voltage of 100 kV. HRTEM was performed by a JEOL JEM-2010 operated at an accelerating voltage of 120 kV. SEM was performed by a Tescan VEGA3 at an accelerating voltage of 20 kV. Raman spectra were recorded by a Renishaw RM2000 spectrometer with 514 nm laser excitation. UV-vis spectra were recorded by a Pgeneral TU-1810 twin-beam spectrometer within a wavelength range of 200–900 nm. FTIR spectra were recorded by a Nicolet 560 spectrometer. XPS spectra were recorded by an Escalab 250 Xi spectrometer.
**S2. UV-vis spectrum of a MoS\(_2\)/DMF solution**

**Fig. S1** UV-vis spectrum of a MoS\(_2\)/DMF solution within a wavelength range of 200–900 nm. The spectrum shows two well-resolved characteristic peaks, *i.e.*, \(A_1\) at 673 nm and \(B_1\) at 613 nm, within 600–700 nm. The two peaks are the first (\(n = 1\)) members of two excitonic Rydberg series that are associated with the direct transition and their energy separation can be ascribed to the spin-orbit splitting of the top of the valence band at the K point [6]. The presence of peaks \(A_1\) and \(B_1\) in the UV/vis spectrum demonstrates the successful exfoliation of MoS\(_2\) into nanosheets in DMF.
Fig. S2  (a) Photograph of a brilliant yellow GO/DMF solution; (b) TEM image of a large-area, highly crumpled GO nanosheet and the corresponding ED pattern (insert).
**S4. UV-vis spectrum of a GO/DMF solution**

**Fig.S3** UV-vis spectrum of a GO/DMF solution within a wavelength range of 200–900 nm. Generally, there should be two characteristic peaks, *i.e.*, a maximum at ~230 nm, presumably due to the π→π* transition of the C–C bonds; a shoulder at ~300 nm, presumably due to the n→π* transition of the C=O bonds [2]. However, the two peaks are invisible because the data within 200–280 nm cannot be collected because DMF has strong absorption in that wavelength range.
**S5. Photograph of a MoS$_2$/GO/PEO system before and after coordination**

![Photograph of a MoS$_2$/GO/PEO system before and after coordination](image)

**Fig. S4** Photograph of a MoS$_2$/GO/PEO (6/3/1) system before (left) and after (middle) coordination, and a MoS$_2$/GO (6/3) system after coordination for comparison (right).
**S6. FTIR spectra of a MoS$_2$/GO/PEO system before and after coordination**

![FTIR spectra](image)

**Fig. S5** FTIR spectra of a MoS$_2$/GO/PEO system (6/3/1) before (green) and after (orange) coordination.

Since the FTIR spectrum of MoS$_2$ is featureless within the wavenumber range of 500–3000 cm$^{-1}$, the characteristic peaks mainly arise from GO. Clearly, before coordination the FTIR spectrum has five characteristic peaks, *i.e.*, C=O bonds (carboxyl/carbonyl) at 1734 cm$^{-1}$, C=C bonds (aromatic ring) at 1647 cm$^{-1}$, C–OH bonds (carboxyl) at 1414 cm$^{-1}$, C–O bonds (epoxide/ether) at 1216 cm$^{-1}$, and C–O bonds (alkoxy) at 1090 cm$^{-1}$. After coordination, the C=O (carboxyl/carbonyl) peak is nearly invisible, and the C–OH (carboxyl) peak increases significantly, indicating the successful coordination between C=O bonds and Cu$^{2+}$ ions. Besides, the C–O (epoxide/ether) peak also disappears, suggesting the possible ring-opening coordination between the C–O bonds and Cu$^{2+}$ ions [7]. This result is in accord with the XPS result shown in Fig. 2.
**S7. Raman spectra of a coordinated MoS$_2$/GO/PEO system before and after reduction**

![Raman spectra graph](image)

**Fig. S6** Raman spectra of a coordinated MoS$_2$/GO/PEO (6/3/1) system before (green) and after (red) reduction. Before reduction, the Raman spectrum of the coordinated MoS$_2$/GO/PEO system has four characteristic peaks. The two weaker ones at 381 and 407 cm$^{-1}$ correspond to the in-plane E$_{2g}$ and out-of-plane A$_{1g}$ vibrations of MoS$_2$, respectively [8]. The two stronger ones at 1351 and 1584 cm$^{-1}$ correspond to the D and G peaks of GO, respectively [9]. After reduction, the D/G intensity ratio increases significantly, suggesting the a decreased average size of the sp$^2$ domains upon the reduction of GO to r-GO. This phenomenon is ascribed by Ruoff et al. to the creation of new graphitic domains after reduction, which are smaller in the size but larger in the number [10].
S8. Photograph of coordinated MoS$_2$/r-GO/PEO systems

Fig. S7  Photograph of three coordinated MoS$_2$/r-GO/PEO systems (7/2/1, 6/3/1 and 5/4/1).
**S9. Photographs of freestanding MoS$_2$/r-GO/PEO (6/3/1) hybrid paper**

**Fig.S8** Photograph of a piece of freestanding MoS$_2$/r-GO/PEO (6/3/1) hybrid paper peeled off the PVDF membrane and tailored arbitrarily by scissors.

**Fig.S9** Photograph of a naked copper foil (left) and a copper foil attached with the tailored freestanding MoS$_2$/r-GO/PEO (6/3/1) hybrid paper (right). Note that there is still scope for improving the electrochemical performance of our film anode by, *e.g.*, rationally reducing its thickness to shorten the diffusion pathways, or trying to further optimise its composition.
Fig. S10. SEM image of a MoS$_2$/GO hybrid directly cross-linked by Cu$^{2+}$ ions (without PEO).
S11. Supplementary references


