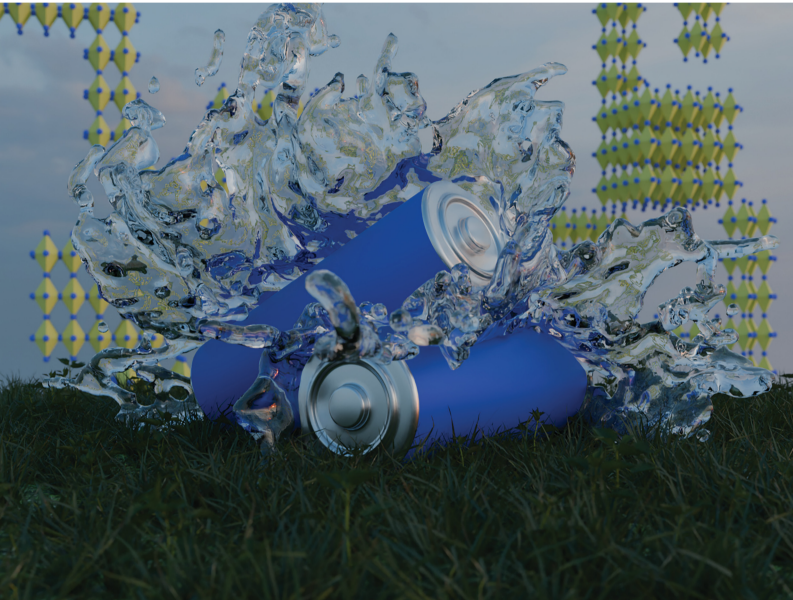
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A nanoengineered vanadium oxide composite as a high-performance anode for aqueous Li-ion hybrid batteries[†]

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Aqueous lithium-ion batteries (LIBs) have received increasing attention as a promising solution for stationary energy storage systems due to their low environmental impact, non-flammability and low cost. Despite recent progress in electrolyte development and cathode manufacturing, the lack of anode materials with high specific capacity presents difficult challenges for a wide range of applications. In this study, we propose a novel synthetic strategy to fabricate a pseudocapacitive V_2O_5 /graphene composite as a highly functional anode material for aqueous LIBs. The designed synthesis combines a fast laser-scribing step with controlled calcination to tune the morphology and oxidation state of the electrochemically active vanadium oxide species while obtaining a highly conductive graphene scaffold. The optimized V₂O₅/graphene anode shows an outstanding specific capacity of 158 mA h g⁻¹ in three-electrode measurements. When the V2O5/graphene anode is paired with an LiMn₂O₄ cathode, the charge storage mechanism of the full cell is revealed to be dominantly surface-controlled, resulting in remarkable rate performance. Specifically, the full cell can reach a specific capacity of 151 and 107 mA h (g anode)⁻¹ at C/6 and 3C, respectively. Moreover, this hybrid battery can achieve a high power density and an energy density of 650 W kg⁻¹ at 15.6 W h kg⁻¹ and 81.5 W h kg⁻¹ at 13.6 W kg⁻¹, respectively, outperforming most aqueous LIBs reported in the literature. This innovative strategy provides a pathway to incorporate pseudocapacitive electrodes for improving aqueous lithium-ion storage systems, enabling safe operation of large-scale energy storage without compromising their electrochemical performance.

New concepts

This work reports an original synthetic approach combining laserscribing and calcination to make a pseudocapacitive V₂O₅/laser-scribed graphene (LSG) composite made from VCl₃/graphene oxide. The laser step generates the 3D graphene scaffold with nanosized vanadium oxide particles. The calcination step converts all remaining vanadium species to V_2O_5 nanoparticles in a controlled manner. In this way, the resulting V_2O_5 nanoparticles in the V2O5-LSG electrode are electrochemically active, small in grain size and easily accessible for charge transport via the LSG structure. The V2O5-LSG composite can be used as an anode in aqueous lithium-ion batteries. Based on previous literature reports, the choice of a suitable transition metal redox couple for an aqueous LIB anode is limited, restricted by low specific capacity. This new synthetic V2O5-LSG anode shows high capacity compared with other aqueous lithium-ion battery anodes. When paired with a LiMn₂O₄ cathode, the full cell displays a predominantly surface-controlled charge storage mechanism that is Faradaic in nature, leading to excellent fast charging properties, and high power and energy density that compare favorably to previously reported aqueous LIBs. Because of its high capacity, fast charging capability and safe operation, the V2O5-LSG||LiMn2O4 system provides a potential solution for future energy storage applications.

Introduction

Aqueous rechargeable batteries have emerged as a promising technology for applications in large-scale energy storage systems due to their safe operation, non-toxicity, and cost-efficiency.¹ Among the storage systems based on various cations such as Na⁺, K⁺, Zn²⁺, Mg²⁺, Al³⁺, *etc.*, aqueous lithium-ion batteries (LIBs) are the most prominent candidates because of the well-developed manufacturing capability for LIB component production. The progress of aqueous LIB research has been significantly advanced by the development of numerous aqueous electrolytes, extending the electrochemical window beyond the conventional 1.23 V limit imposed by the electrolytic decomposition of water.^{2–5} A "water-in-salt" electrolyte was therefore obtained by dissolving lithium bis(trifluoromethane sulfonyl)-imide (LiTFSI) at extremely high concentrations (molality > 20 M)

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in water, which can widen the electrochemical window to >3 V. LiTFSI was chosen as the salt because of its high solubility in water (>20 M at 25 °C) and high stability against hydrolysis. In such a super-concentrated electrolyte, the decomposition of salt anion occurs preferentially on the anode before hydrogen evolution occurs, leading to the formation of a dense solid electrolyte interphase (SEI) primarily consisting of LiF to solve the hydrogen and oxygen evolution problem.⁵ While the Li/Li⁺ intercalation potential of commercially available LIB cathodes including LiCoO₂, LiFePO₄, and LiMn₂O₄ conveniently falls below the upper potential limit of the operational window of aqueous electrolytes, carbon-based anodes that are widely used in organic electrolytes are not compatible with the aqueous system due to their low Li/Li⁺ intercalation metal-based anodes must be employed in aqueous LIBs.

Restricted by the relatively high lower potential limit of aqueous lithium-ion electrolytes, the choice of suitable transition metal redox couples for an aqueous LIB anode is limited. Most reported aqueous LIBs use an intercalation-type Mo₆S₈ or TiO₂ anode. Although high average full-cell voltage can be achieved, the energy density is restricted by their low specific capacity, since a high capacity ratio between the negative electrode (anode) and positive electrode (cathode), known as the n:p ratio, is required to compensate for the initial loss of anode capacity by reactions with the electrolyte.⁵ Therefore, there is an urgent need for anode materials that exhibit high specific capacity. Vanadium oxide anodes have been reported to show enhanced capacity in aqueous LIBs; however, the added capacity is not sufficient to account for the lower average voltage owing to their unfavorable redox potential, leading to disadvantageous energy density.¹²⁻¹⁵ Moreover, aqueous LIBs have great potential for fast charging applications due to the enhanced diffusion in an aqueous electrolyte, which can be further improved by integrating electrodes with multidimensional diffusion pathways or with redox reactions confined to the surface. The spinel LiMn₂O₄ cathode allows threedimensional Li migration, and V2O5, as an intrinsically intercalation-type material, can show extrinsic pseudocapacitive behavior when nanosized.¹⁶⁻¹⁹ Therefore, the combination of these three components is expected to deliver a hybrid battery with excellent fast charging capability.

In this work, we report a novel V_2O_5 /laser-scribed graphene (LSG) composite material in which V_2O_5 nanoparticles are uniformly anchored on a conductive graphene nanosheet scaffold. The V_2O_5 -LSG composite is made from VCl₃/graphite oxide *via* an originally designed two-step laser scribing/calcination approach where the former results in an expanded LSG

network as well as nanosized vanadium particles and the latter further converts all vanadium species to V2O5 nanoparticles in a controlled manner. In this way, the resulting V₂O₅ nanoparticles in the V₂O₅-LSG electrode are electrochemically active, small in grain size and easily accessible for charge transport via the LSG structure. This V₂O₅-LSG electrode shows pseudocapacitive features and remarkable rate capability in three-electrode measurements. When the V2O5-LSG anode is paired with a LiMn₂O₄ cathode, the hybrid battery exhibits predominantly surface-controlled charge storage behavior, leading to excellent fast charging capability. The full-cell specific capacity can reach 151 mA h (g anode)⁻¹ giving high power and energy density, 650 W kg⁻¹ at 15.6 W h kg⁻¹ and 81.5 W h kg⁻¹ at 13.6 W kg⁻¹, which outperform those of the majority of previously reported aqueous LIBs. Moreover, the effects of the calcination time were studied by transmission electron microscopy (TEM) and X-ray photoelectron spectroscopy (XPS), revealing how the chemical position and structural morphology of the electrochemically active vanadium species can be effectively optimized.

Results and discussion

The synthesis of V₂O₅-laser scribed graphene (LSG) was performed to adapt a pseudocapacitive material for battery chemistry. As shown in Fig. 1, the VCl₃ solution was added to a graphite oxide (GO) dispersion to ensure uniform mixing of vanadium ions and GO sheets. The vanadium cations are believed to act as spacers to avoid aggregation of GO layers. As reported in our prior work, laser-scribing of a drop-cast VCl₃/ GO film can simultaneously lead to the formation of low-valent vanadium oxides (VO_x , predominantly V_2O_3 and VO_2) and an expanded reduced graphene oxide (rGO) network due to the rapid release of gases, denoted here as VO_x-LSG.^{20,21} Although laser-scribing alone can fabricate thin-film electrodes for supercapacitor systems that only rely on surface charge storage processes, the electrodes usually suffer from low areal mass loading^{22,23} and incompatibility with battery-type electrochemistry, which require more uniformly formed electrodes to carry out ion transport deeper beneath the surface. Therefore, after the transient heat exposure induced by a CO₂ laser, the VO_x-LSG was exposed to a much longer thermal treatment in air, resulting in the oxidation of lower-valent VO_r with an intrinsic tunnel-like structure and one-dimensional charge transfer pathways to a layered V₂O₅ structure where twodimensional diffusion is enabled.²⁴ By choosing a sufficiently

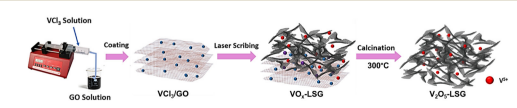


Fig. 1 Schematic illustration of V₂O₅-LSG synthesis *via* laser scribing and calcination.

low temperature, thermal decomposition of graphene to CO is prevented and the integrity of the LSG structure is preserved. In the end, the transformation from any unconverted VCl₃/GO and VO_x-LSG intermediate into a V₂O₅-LSG composite was completed.

The structural features of the as-synthesized V₂O₅–LSG composite were investigated by electron microscopy (SEM) and transmission electron microscopy (TEM). In Fig. 2a, randomly stacked rGO flakes with visible wrinkles are observed in the SEM image, suggesting the successful formation of the conductive LSG network. Under high magnification, Fig. 2b shows that the LSG sheets are evenly and fully coated with V₂O₅ nanoparticles. From the TEM images (Fig. 2c), it can be observed that nanoparticles are uniformly distributed on the graphene sheets and are ~40–50 nm in size. Fig. S15 (ESI†) shows the size distribution of VO_x particles based on Fig. 2c and the vast majority of the sizes are ~40–50 nm. The SEM images of the precursor VCl₃/GO and the intermediate VO_x–LSG

are displayed in Fig. S2 (ESI[†]). Fig. S2a and b (ESI[†]) show flat GO sheets closely stacked together mixed with an aggregated network of VCl₃ while the VO_x–LSG in Fig. S2c and d (ESI[†]) has expanded spacing between the rGO sheets and no sign of agglomerates of VO_x, demonstrating that the laser step simultaneously achieves reduction of GO and nanosizing of the vanadium species. The morphological evidence indicates that the interconnected and porous LSG structures are preserved and the V₂O₅ particle size remained within the same order of magnitude after calcination.

The compositions of the synthesized V₂O₅–LSG composites were studied by X-ray powder diffraction (XRD) and Raman spectroscopy. Fig. 2d presents the diffraction pattern of the V₂O₅–LSG composite and it matches that of orthorhombic V₂O₅, confirming the formation of V₂O₅ from the lower-valent VCl₃ precursor and the VO_x intermediate. The most prominent peaks at 20.4°, 26.3° and 31.2° correspond to the (001), (110) and (310) planes, respectively. The rest of the peaks are VO₂ and

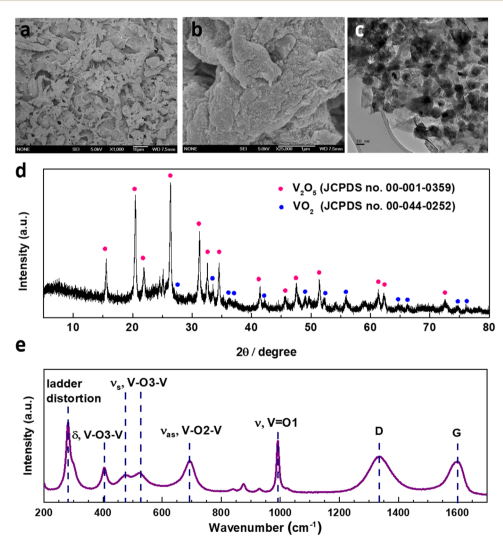


Fig. 2 Characterization of the as-synthesized V_2O_5 -LSG composite (calcination time = 1 h). (a) Low- and (b) High-magnification SEM images of the V_2O_5 -LSG composite. (c) A TEM image showing the V_2O_5 particles on the rGO sheets. (d) XRD pattern of V_2O_5 -LSG matching V_2O_5 (JCPDS no. 00-001-0359) and VO_2 (JCPDS no. 00-044-0252). (e) Raman spectra of the composite showing V_2O_5 and graphene features.

match with JCPDS no. 00-044-0252. Fig. S3a and b (ESI⁺) show the XRD patterns of VCl₃/GO and VO_x-LSG, respectively. The former resembles the patterns of graphite and graphite oxide with no signs of any crystalline vanadium oxide species. The latter shows peaks attributed to V₂O₃, VO₂ and mixedvalent vanadium oxides, indicating that the laser-scribing step involves the oxidation from VCl₃ to the multivalent VO_x and the following calcination completes further oxidation to V₂O₅. The successful synthesis of V2O5-LSG was confirmed using Raman spectra, which show all the V₂O₅ vibrational features as well as the D and G bands of graphene (Fig. 2e). O1, O2 and O3 denote the out-of-plane terminal, in-plane terminal and in-plane bridging oxygen atoms in V_2O_5 , and all the peaks are labeled with the corresponding vibrational modes.²⁵ Furthermore, the complete oxidation of VCl₃ to V₂O₅ is confirmed by the absence of both Cl 2s and Cl 2p peaks in the XPS survey spectrum of V_2O_5 -LSG (Fig. S1, ESI^{\dagger}). Furthermore, since the calcination temperature was chosen to be below 400 °C, above which LSG is vulnerable to thermal oxidation to CO, $\sim 10\%$ of the final V₂O₅ composite is LSG by weight, as signified by the sharp decrease around 400 °C in the thermogravimetric analysis (TGA), as shown in Fig. S4 (ESI[†]). Therefore, we choose a temperature lower than 400 °C to prevent the thermal decomposition of graphene to CO and preserve the integrity of the LSG structure. We made the assumption that at 300 °C the conversion to V₂O₅ can be completed without risking the loss of the rGO scaffold. Overall, all characterization results indicate that the post-laserscribing calcination step ensures full reaction converting VCl₃ to V₂O₅ without compromising the conductive LSG scaffold, as illustrated in Fig. 1.

The electrochemical features of the V₂O₅–LSG composite electrodes were first studied using a three-electrode setup in the potential window 0 to -1.4 V vs. Ag/AgCl with 21 M lithium bis(trifluoromethane)sulfonimide (LiTFSI) as the electrolyte. As shown in Fig. 3a, the CV curves possess broad peaks with small peak-to-peak separations, which is a feature of fast Faradaic charge storage processes and is typical for a redoxactive pseudocapacitive material. The voltage profile in Fig. 3b shows two sloping regions with drastically different gradients. From -1.1 V to -0.5 V vs. Ag/AgCl, there is a sloping plateau and the capacity contribution gradually decreases with increasing current density, indicting the rate-dependent battery-type behavior. This is consistent with CV measurements since the most pronounced redox peaks are also observed in the same region. On the other hand, a sloping voltage is always observed between -0.5 V and 0 V *vs.* Ag/AgCl and the gradient remains constant with changing current density, signifying capacitive behavior. As described in Fig. 3c, the V₂O₅–LSG composite electrode demonstrated an excellent rate capability in 21 M LiTSFI. At 0.1, 0.2, 0.5 and 1 A g⁻¹, the average discharge capacity can reach 158, 145, 131 and 119 mA h g⁻¹, respectively, based on the total mass of the V₂O₅–LSG composite. The LSG contribution is limited to ~11 mA h g⁻¹ based on the three-electrode test shown in Fig. S14 (ESI[†]). The specific capacity per VO_x is calculated as follows:

Specific capacity per VO_x

$$= \frac{(\text{total capacity} - \text{LSG spec.capacity} \times \text{LSG mass})}{\text{VO}_x \text{ mass}}$$
(1)
$$= \frac{158 - 11}{0.839} = 175.2 \text{ mA h g}^{-1}$$

Impressively, 75% of the maximum capacity was retained upon a 10-fold rate increase from 0.1 to 1 A g⁻¹. Additionally, ~7% capacity loss was observed in the first cycle, which is speculated to be caused by irreversible formation of LiF from the electrode reaction with LiTSFI.⁵ When the rate is returned to 0.1 A g⁻¹ in the end, ~95% of the initial capacity was recovered after accounting for the loss from LiF formation. The outstanding rate performance of the V₂O₅-LSG electrode provides a promising foundation for its fast-charging applications in full cells.

To evaluate the electrochemical performance of the V₂O₅–LSG composite in a more practical setting, it was tested as the anode in full cells and paired with an intercalation-type cathode LiMn₂O₄, as illustrated in Fig. 4a. Fig. 4b shows the CV curves of the V₂O₅–LSG anode and the LiMn₂O₄ cathode in 21 M LiTFSI electrolyte. The 21 M LiTFSI electrolyte has a wider electrochemical window with its hydrogen and oxygen evolution potentials extended to -0.9 V and 2 V, respectively, compared with 5 M LiTFSI which only has an electrochemical window of 1.7 V (Fig. S16a, ESI[†]),

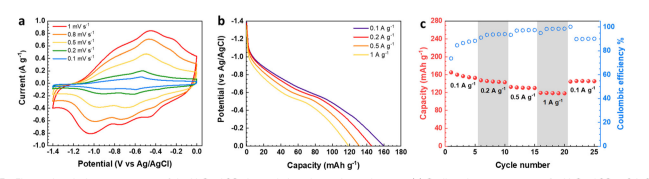


Fig. 3 Electrochemical measurements of the V_2O_5 -LSG electrode in a three-electrode setup. (a) Cyclic voltammetry curves for V_2O_5 -LSG at 0.1, 0.2, 0.5, 0.8, and 1 mV s⁻¹. (b) Capacity-voltage profiles at 0.1, 0.2, 0.5, and 1 A g⁻¹. (c) Rate performance and corresponding Coulombic efficiency of V_2O_5 -LSG cycled at different current densities.

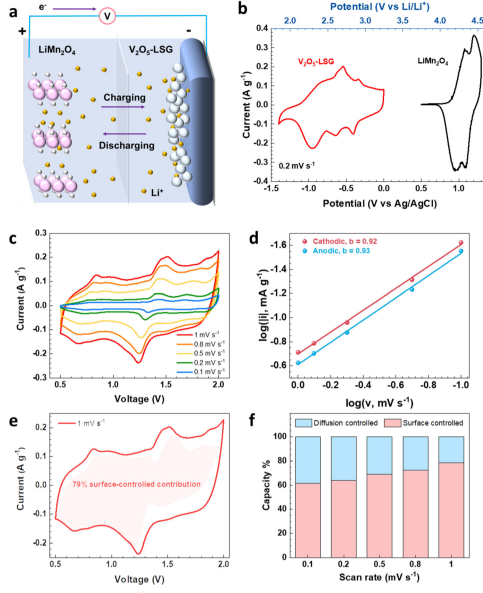


Fig. 4 Electrochemical analysis of the $V_2O_5-LSG||LiMn_2O_4$ cells. (a) Illustration of Li-ion storage in the hybrid system based on a pseudocapacitive anode and an intercalation-type cathode. (b) Potential windows of V_2O_5-LSG and $LiMn_2O_4$ in a 21 M LiTFSI electrolyte. (c) Cyclic voltammetry curves of a $V_2O_5-LSG||LiMn_2O_4$ coin cell at 0.1, 0.2, 0.5, 0.8, and 1 mV s⁻¹. (d) *b*-values of the most pronounced pair of peaks in the CV. (e) Current contribution by surface-controlled processes at 1 mV s⁻¹. (f) Capacity contribution by surface-controlled and diffusion-controlled processes at a range of scan rates.

indicating a sufficiently wider electrochemical window of ~3 V that ensures stable operation of both the V₂O₅–LSG anode and the LMO cathode. The anode is electrochemically active between 1.8 V and 3.2 V vs. Li/Li⁺, which covers the potential range of common aqueous lithium-ion battery anodes such as Mo₆S₈ and TiO₂.^{4,26} The LMO cathode is electrochemically active between 3.8 V and 4.6 V vs. Li/Li⁺, which enables a wider voltage window. Another cathode, LiFePO₄, was tried to pair with the anode. However, LFP has a lower potential range compared to LMO (Fig. S17a, ESI⁺). Therefore, LMO with a higher potential was used enabling a wider voltage window and a higher capacity (Fig. S17b, ESI⁺).

The discharge capacity of the $LiMn_2O_4$ cathode was measured to be 112 mA h g⁻¹ (Fig. S5, ESI⁺), which is 29% lower

than that of the V_2O_5 -LSG anode. Conventional LIB anodes like TiO₂⁴ or Mo₆S₈,⁵ which have similar specific capacities of 125 mA h g⁻¹ and 118 mA h g⁻¹, respectively, like LiMn₂O₄, often require doubling the mass of the anode to obtain a suitable n:p ratio in order to compensate for the anode capacity loss due to solid–electrolyte interphase formation;⁶ however, since the V₂O₅-LSG anode capacity is 41% higher than that of LiMn₂O₄, an anode : cathode mass ratio of 1 was used in all full cell testing. In addition, the cell capacity is based on the total mass of the anode and cathode. We have carried out experiments with higher and lower cathode masses (Fig. S13, ESI†) and found that when the mass ratio is 1:1, the full cell has the highest capacity (mA h g_{total mass}⁻¹). When the mass ratio

V₂O₅-LSG: LiMn₂O₄ is 1:2, the capacity decreases. The reason is that when increasing the cathode mass, while the capacity increases, the total mass also increases, but the ratio of capacity/total mass decreases because the total mass increases more. When the mass ratio V₂O₅-LSG: LiMn₂O₄ is 2:1, the capacity also decreases. The reason is the same. When increasing the anode mass, the capacity increases; however, the total mass also increases, but the ratio of capacity/total mass decreases because the total mass increases more. As shown in Fig. 4c, between 0.5 and 2 V, the CV curves of the V₂O₅-LSG||LiMn₂O₄ cell adopt a distorted rectangular shape with prominent redox peaks, which is typical for a hybrid energy storage system.

In order to investigate the kinetics of the charge storage processes in the V₂O₅-LSG||LiMn₂O₄ cell, the sharpest pair of peaks that appear between 1.2 and 1.6 V were analyzed. The peak current densities (*i*) and the scan rates (ν) generally obey the following power law:^{27,28}

$$i = av^b \tag{2}$$

By plotting log(*i*) of each peak against log(ν), the *b*-value of the peak can be determined, and the common consensus is that *b*-values of 0.5 and 1 indicate battery-type and capacitortype behavior, respectively. Fig. 4d illustrates that the anodic and cathodic peaks, respectively, have *b*-values of 0.93 and 0.92 (with $R^2 = 0.998$), suggesting that the charge storage mechanism of the V₂O₅–LSG||LiMn₂O₄ system consists of both battery-type and capacitor-type processes that is dominated by the latter. Furthermore, to gain more quantitative insights into the charge storage behavior and decouple the capacity contribution from the diffusion-controlled battery-type and the surface-controlled capacitor-type processes, a second kinetic analysis was carried out based on the following relationship:

$$i = k_1 v + k_2 v^{\frac{1}{2}} \tag{3}$$

The surface-controlled component (k_1v) and the diffusioncontrolled component $(k_2 v^{\frac{1}{2}})$ are proportional to the scan rate and the square root of the scan rate, respectively. Fig. 4e shows an example where the current contributions from the two types of processes were quantified based on the CV data of the system at 1 mV s⁻¹. It is clear that the lower-voltage region (0.5–1.2 V) is more predominantly governed by surface-controlled processes, whereas the diffusion-controlled current starts to significantly grow going into the higher-voltage region (1.2-2 V). This is consistent with the observation in Fig. 3b, in which the V_2O_5 -LSG anode also presents more battery-type behavior toward higher state-of-charge. This analysis was also carried out using CV data at 0.1, 0.2, 0.5 and 0.8 mV s^{-1} and the corresponding capacity quantification is summarized in Fig. 4f. The capacity contributed by the diffusion-controlled processes decreased from 79% at 1 mV s⁻¹ to 61% at 0.1 mV s⁻¹. This indicates that the charge storage mechanism of the V2O5-LSG||LiMn2O4 system is mainly governed by capacitive processes even at very slow scan rates although the contribution gap between the processes gradually lessens from 1 to 0.1 mV s⁻¹. This capacity

dominated by diffusion-controlled contributions of V_2O_5 -LSG||LiMn₂O₄ gives an excellent foundation for its great electrochemical performance at high rates.

The electrochemical performance of the V₂O₅-LSG||LiMn₂O₄ system was assessed in coin cell formats. First, a control experiment was carried out to eliminate any capacity contribution from the graphite paper substrate, and Fig. S6 (ESI[†]) shows that the substrate has a negligible capacity when paired with a $LiMn_2O_4$ cathode. The V₂O₅-LSG||LiMn₂O₄ cell was cycled at C/6, C/2, 1C, 3C and C/6 rates as shown in Fig. 5a. The average discharge capacity reached 151, 137, 123 and 107 mA h (g anode) $^{-1}$, or 75, 69, 61 and 53 mA h g^{-1} based on the total electrode mass, at C/6, C/2, 1C, and 3C, respectively. With a 18-fold rate increase from C/6 to 3C, 71% capacity was retained, signifying outstanding fast charging capability. Again, the same initial decrease in capacity is observed, leading to 91% capacity retention when the rate is reversed to C/6, as seen in Fig. 3c. The voltage profiles of the V₂O₅-LSG||LiMn₂O₄ cell consist of a small plateau around 1.5 V and a voltage slope in the 0.5-1.4 V range, indicating that the capacity is predominantly from surface-controlled processes, as illustrated in Fig. 5b. The plateau is a feature of diffusioncontrolled Faradaic processes, and its gradient increases with increasing rate, suggesting that the capacity contribution from battery-type behavior also falls with the increasing rate. This trend corroborates with the kinetic analysis results presented in Fig. 4f, and the plateau voltage matches the peak positions in the CV curves (Fig. 4c). Faster conditions have been tested with 6C (Fig. S18, ESI[†]). With a 6-fold rate increase from 1C to 6C, 65% of the capacity was retained, signifying outstanding fast charging capability. When the rate is returned to 1C, $\sim 95\%$ of the initial capacity was recovered, confirming the fast charging capacity.

The V₂O₅-LSG||LiMn₂O₄ system was tested for long-term cycling at a 1C rate (Fig. 5c). 86% and 75% of the initial capacity were retained after 50 and 100 cycles, respectively. The SEM images of the V2O5-LSG electrodes before and after 100 cycles demonstrated that there are no significant morphological changes and the LSG network remains intact after 100 cycles (Fig. S11, ESI[†]). As illustrated in Fig. 5d, the capacity fading is associated with the disappearance of the voltage plateaus around 0.8 and 1.5 V, indicating reduced contribution from diffusion-controlled Faradaic processes. Moreover, the LSG||LiMn₂O₄ cell showed excellent long-term cycling stability at a high rate of 5C (Fig. 5e), reaching 80% and 62% capacity retention after 330 and 800 cycles, respectively. As shown in Fig. 5f, the individual plateaus are less observable in the charge and discharge curves during 5C cycling, displaying predominantly pseudocapacitive behavior. Furthermore, when a pair of V_2O_5 -LSG||LiMn₂O₄ cells were connected in series, the battery pack can light up not only red, but also blue LED lights; impressively, the fully charged battery pack can power the LED for more than 24 h (Fig. S7, ESI⁺). Additionally, several control experiments were carried out to confirm the source of the great electrochemical performance and to evaluate the designed synthetic strategy. Fig. S8 (ESI[†]) compares the electrochemical properties of a physical mixture V₂O₅ + rGO to those of the nanoengineered V₂O₅-LSG, and the former shows

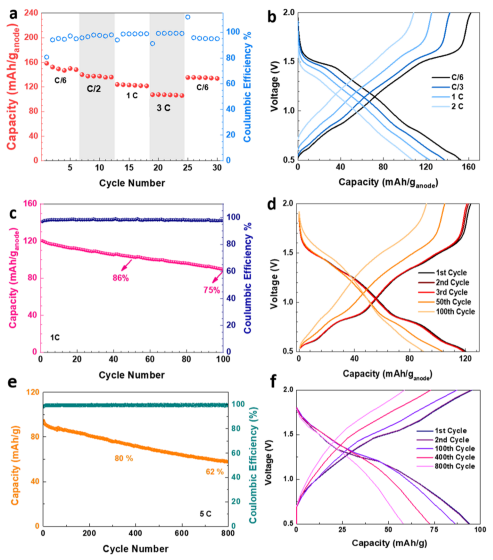


Fig. 5 Electrochemical performance of the V_2O_5 -LSG||LiMn₂O₄ coin cells. (a) Rate performance and corresponding Coulombic efficiency of a V_2O_5 -LSG||LiMn₂O₄ cell cycled at C/6, C/2, 1C, 3C and C/6. (b) Capacity-voltage profiles at different rates. (c) Capacity and Coulombic efficiency of a V_2O_5 -LSG||LiMn₂O₄ cell over 1C cycling. (d) Capacity-voltage profiles of the 1st, 2nd, 3rd, 50th, and 100th cycles during 1C cycling. (e) Capacity and Coulombic efficiency of a V_2O_5 -LSG||LiMn₂O₄ cell over 5C cycling. (f) Capacity-voltage profiles of the 1st, 2nd, 100th, 400th and 800th cycles during 5C cycling.

poor rate capability and cyclability due to the lack of a conductive scaffold and V_2O_5 nanosizing. As illustrated in the electrochemical impedance spectroscopy (EIS) spectra, the system with V_2O_5 -LSG possesses dramatically reduced charge transfer resistance compared to the physical mixture (Fig. S8c, ESI†). The V_2O_5 -LSG composite electrode fabricated following the synthesis in Fig. 1 is also compared to a sample that only underwent one of either the laser-scribing or calcination steps (Fig. S9, ESI†). Both the calcination only and laser only samples experienced a sharp capacity decrease at the start of cycling and showed inferior rate capability, confirming the necessity of both steps. The poor performance of the laser only sample can be attributed to the incomplete oxidation and a large amount of the remaining VCl₃ precursor (Fig. S1, ESI†), which is susceptible to dissolution into the electrolyte. As shown in Fig. S9c (ESI[†]), the Nyquist plots of the systems with V₂O₅–LSG and VO_x–LSG are similar with significantly lower resistance than the VCl₃/GO precursor mixture, confirming that a highly conductive network is formed upon laser scribing. As shown in the SEM images of the calcination only electrode (Fig. S10a, ESI[†]), without the laser-scribing step, the graphene sheets are less expanded in comparison to the porous 3D LSG structure (Fig. 2a). In addition, large V₂O₅ clusters of ~2 µm are observed, which are much larger than the nanoparticles in the V₂O₅–LSG composite, leading to a reduced active surface area and poor rate capability (Fig. S10b and Fig. 2c, ESI[†]). Overall, the meticulously designed V₂O₅–LSG electrode exhibits a high specific capacity and excellent fast-charging properties due to its predominantly surfacecontrolled charge storage mechanism that is triggered by the V₂O₅-nanoparticle-on-LSG-scaffold structure.

Furthermore, the calcination time in the synthesis to convert VO_x-LSG to V₂O₅-LSG (Fig. 1) was varied to investigate its effects on the resulting composite. The TEM images of V₂O₅-LSG that underwent post-laser calcination for 0.5 h, 1 h and 2 h are shown in Fig. 6a–c. In all three images, VO_x/V_2O_5 is distributed on graphene sheets and its morphology changes significantly with increasing calcination time. With only 0.5 h post-laser calcination, the VO_x/V₂O₅ clusters adopt undefined shapes and are uneven in size, resembling the morphology of spinodal decomposition products (Fig. 6a).²⁹ As the calcination time increased to 1 h, defined V₂O₅ nanoparticles formed with an average size of ~ 50 nm (Fig. 6b and Fig. 2c). The nucleation process continued from 1 h to 2 h, inducing much larger agglomeration of V₂O₅ particles by Ostwald ripening (Fig. 6c).³⁰ X-ray photoelectron spectroscopy (XPS) was used to elucidate the changes in the chemical composition of the VO_x/V₂O₅-LSG composites and the deconvoluted V 2p regions are shown in Fig. 6d-f and Fig. S12 (ESI[†]). In Fig. 6d, the peak at 516.6 eV represents V4+ and accounts for 58% of all V present, indicating that the major oxidation state of VO_x is 4+ after 0.5 h post-laser calcination and the oxidation process is incomplete. As depicted in Fig. 6e and f, after 1 h and 2 h of calcination, the V⁵⁺ peak at 517.8 eV becomes the dominant peak in the spectrum and makes up ~80% of all V present, suggesting that V⁵⁺ has overtaken V⁴⁺ as the main vanadium species present. Fig. S12 (ESI⁺) shows the V 2p sub-spectrum of the VO_x–LSG intermediate that has never undergone any calcination, and it shows the presence of vanadium oxidation states +2 to +5, indicating incomplete oxidation of the VCl₃ precursor without the calcination step. Thus, TEM images and XPS V 2p spectra suggest that it takes more than 0.5 h for the oxidation from VO_x to V₂O₅ to reach completion and the optimal particle size is obtained after 1 h of calcination.

The electrochemical behavior of the composites with various calcination times was also studied in full cells when they are paired with LiMn_2O_4 cathodes. Their average discharge capacities of cycle 25–30 (where the low-current capacities were stabilized) are compared in Fig. 6g. The cells with 0.5 h, 1 h and 2 h have specific capacity of 72, 135 and 52 mA h g⁻¹, respectively. As shown by TEM and XPS data, on the one hand, with shorter calcination time, the conversion to V₂O₅–LSG is

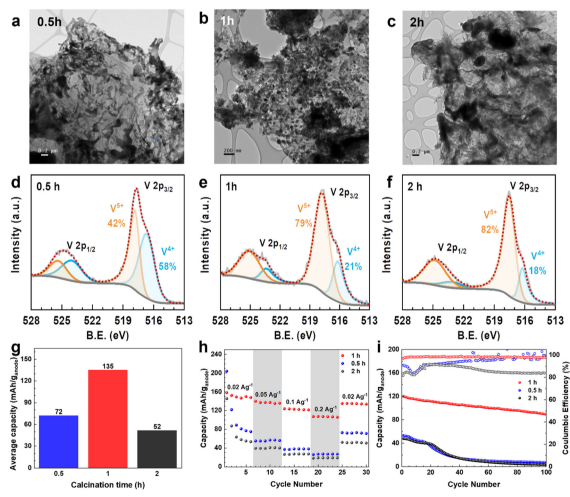


Fig. 6 Effects of calcination time on the V_2O_5 -LSG composite. TEM images of the eV_2O_5 -LSG composite calcined for (a) 0.5 h, (b) 1 h, and (c) 2 h. XPS V2p spectra of the V_2O_5 -LSG composite calcined for (d) 0.5 h, (e) 1 h, and (f) 2 h. (g) Specific capacity comparison between samples synthesized with 0.5, 1, and 2 h of calcination. (h) Rate performance of V_2O_5 -LSG||LiMn₂O₄ cells with different calcination times cycled at C/6, C/2, 1C, 3C and C/6. (i) Capacity and Coulombic efficiency of V_2O_5 -LSG||LiMn₂O₄ cells with different calcination times over cycling at 0.1 A g⁻¹.

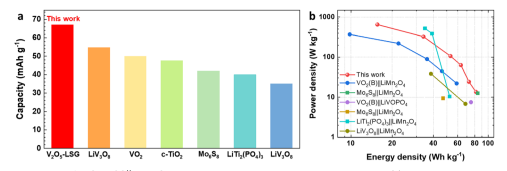


Fig. 7 Performance comparison of $V_2O_5-LSG||LiMn_2O_4$ to previously reported systems in the literature. (a) A capacity comparison based on the total electrode mass of $V_2O_5-LSG||LiMn_2O_4$ to those of other aqueous Li-ion batteries reported in the literature.^{5,11,30-33} (b) A Ragone plot comparing power and energy densities based on the total electrode mass of $V_2O_5-LSG||LiMn_2O_4$ to those of other aqueous Li-ion batteries reported in the literature.

incomplete; on the other hand, the V₂O₅ nanoparticles become aggregated clusters with 2 h of calcination making a large proportion inaccessible near the core. Either way, this leads to a low proportion of the electrochemically active V_2O_5 in the resulting composites, reducing the specific capacity, suggesting 1 h as the optimal calcination duration. Fig. 6h shows the rate performance of full cells with these VO_x/V₂O₅-LSG anodes. Both cells with anodes that underwent calcination for 0.5 h and 2 h show sharp capacity fading during the first 5 cycles at a low current density of 0.02 Ag^{-1} , suggesting possible structural instability caused by incomplete formation of V₂O₅ structures or pulverization accompanied by a large cluster size. When the current density was increased 10-fold, the cells with 0.5 h and 2 h calcination anodes can only reach a capacity retention of 35% and 37%, respectively, whereas 71% of capacity was retained for the anode with 1 h calcination time.³¹

The cycling stabilities of cells with anodes of different calcination times were also investigated (Fig. 6i). The capacity of cells with anodes of 0.5 h and 2 h calcination fell below 26% after only 40 cycles, while the cell with the 1 h calcination anode can retain 86% of its capacity after the same number of cycles. In addition, the Coulombic efficiencies of cells with anodes that underwent 0.5 h and 2 h calcination are low (~80%) during the first 20 cycles. For the 0.5 h calcination anode, this phenomenon can be attributed to the dissolution of incompletely reacted VO_x in the electrolyte. For the 2 h calcination anode, the Coulombic efficiency remained low throughout the cycling, which is likely caused by Li⁺ trapped deep into the bulk of the agglomerated V₂O₅ particles.

Finally, the electrochemical performance of the V_2O_5 -LSG||LiMn₂O₄ system is compared to those of other lithiumion batteries (LIBs) previously reported in the literature. Fig. 7a compares the full-cell specific capacity based on the total mass of both anode and cathode compositions when other aqueous LIB anodes are paired with the LiMn₂O₄ cathode.^{5,13,32-35} The V₂O₅-LSG||LiMn₂O₄ system gives the highest specific capacity of 67 mA h g⁻¹ (after an initial capacity loss is accounted for) compared to other Ti, Mo and V-based anodes. This competitive advantage is attributed to the high capacity of the designed binder-free V₂O₅-LSG anode. The Ragone plot (Fig. 7b) compares the energy and power density of the V₂O₅-LSG||LiMn₂O₄ cell to those of the aqueous LIBs reported in the literature.^{5,13,15,32,33,35,36} The V₂O₅–LSG||LiMn₂O₄ system can reach a high power density of 650 W kg⁻¹ at 15.6 W h kg⁻¹ and a high energy density of 81.5 W h kg⁻¹ at 13.6 W kg⁻¹, outperforming the majority of prior aqueous LIBs. The detailed calculation is shown in Table S1 (ESI†). The outstanding power density can be attributed to the predominantly surface-controlled charge storage as illustrated in Fig. 4f. Notably, although V-based anodes have a relatively less negative redox potential that leads to lower average full-cell voltages in comparison to Ti and Mo-based anodes, the energy density of the V₂O₅–LSG||LiMn₂O₄ cell is actually comparable to if not higher than those of other aqueous LIBs due to its remarkably high anode capacity. Thus, the V₂O₅–LSG||LiMn₂O₄ system exhibits tunable power and energy density and is a promising candidate for fast-charging and safe energy storage applications.

Conclusions

In summary, a pseudocapacitive composite anode for aqueous lithium-ion storage in which V2O5 nanoparticles are distributed on a conductive graphene scaffold is reported for application in aqueous LIBs. The V2O5-LSG composite was synthesized from VCl₃/GO precursors by a two-step laser-scribing/calcination approach, resulting in both an expanded LSG structure and fully oxidized V_2O_5 particles that are ~50 nm in size. The V_2O_5 -LSG anode can achieve an outstanding average specific capacity of 158 mA h g^{-1} and excellent rate capability as shown in threeelectrode measurements. When paired with the intercalation-type cathode LiMn₂O₄ cathode, the full cell displays a predominantly surface-controlled charge storage mechanism that is faradaic in nature, leading to excellent fast charging properties. The V2O5-LSG||LiMn₂O₄ system shows an average specific capacity of 151 and 107 mA h (g anode)⁻¹ at C/6 and 3C. Furthermore, the full-cell specific capacity as well as the power and energy density of this hybrid battery can reach 67 mA h g⁻¹, 650 W kg⁻¹ $(at 15.6 \text{ W h kg}^{-1})$ and 81.5 W h kg $^{-1}$ at (13.6 W kg $^{-1})$ respectively, all of which are favorable compared to those of previously reported aqueous LIBs. Thanks to its high capacity, fast charging capability and safe operation, the V2O5-LSG||LiMn2O4 system provides a potential solution for future energy storage applications.

Conflicts of interest

The authors declare no conflicts of interest.

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