Universal roles of hydrogen in electrochemical performance of graphene: high rate capacity and atomistic origins

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Atomic hydrogen exists ubiquitously in graphene materials made by chemical methods. Yet determining the effect of hydrogen on the electrochemical performance of graphene remains a significant challenge. Here we report the experimental observations of high rate capacity in hydrogen-treated 3-dimensional (3D) graphene nanofoam electrodes for lithium ion batteries. Structural and electronic characterization suggests that defect sites and hydrogen play synergistic roles in disrupting $sp^2$ graphene to facilitate fast lithium transport and reversible surface binding, as evidenced by the fast charge-transfer kinetics and increased capacitive contribution in hydrogen-treated 3D graphene. In concert with experiments, multiscale calculations reveal that defect complexes in graphene are prerequisite for low-temperature hydrogenation, and that the hydrogenation of defective or functionalized sites at strained domain boundaries plays a beneficial role in improving rate capacity by opening gaps to facilitate easier Li penetration. Additional reversible capacity is provided by enhanced lithium binding near hydrogen-terminated edge sites. These findings provide qualitative insights in helping the design of graphene-based materials for high-power electrodes.

The commercial applications of graphene materials for energy storage devices, including lithium ion batteries (LIBs) and supercapacitors, hinge critically on our ability to produce these materials in large quantities and at low-cost. The wet chemistry approach (i.e., liquid phase exfoliation and reduction of graphene oxide) can produce graphene in tons and holds much promise. However, impurities—especially atomic hydrogen—exist ubiquitously in graphene materials made by chemical methods, and their effect on lithium storage capability remains little understood despite some rather important roles that these cross contaminants could play. Theory and experiments have suggested that hydrogen adsorbents impact the electronic structures of graphene and are expected to play a significant role in influencing the lithium storage capacity when applied as anodes in LIBs. Nonetheless, the conclusive evidence for this behaviour has not been forthcoming.

Earlier studies on various carbon materials (e.g., most are soft carbon from pyrolysis of organic precursors) generally pointed at an increased lithium storage capacity with increasing hydrogen content. The exact mechanisms underpinning this empirically observed behaviour remain a subject of ongoing debate and it is unclear whether this phenomenon also occurs for graphene. A main challenge has been to control the hydrogen content and location in graphene materials, a subject that is also of great interest to hydrogen storage applications. Control experiments on graphene single sheets, on the other hand,
Table 1. Processing conditions of graphene nanofoams (GNFs) samples used in this study.

| Sample ID     | Processing conditions                                                                 |
|---------------|----------------------------------------------------------------------------------------|
| GNF-1050C     | Pyrolyzed at 1050°C in N₂ (99.99+%), 4hrs, reference sample                           |
| GNF-1050C-H   | GNF-1050C sample further heat treated at 400°C, 100-sccm flow of 4at% H₂/Ar, 24hrs    |
| GNF-1050C-H₂  | GNF-1050C sample further heat treated at 400°C, 100-sccm flow of 100% H₂, 4hrs        |
| GNF-1600C     | GNF-1050C sample further heat treated at 1600°C, Ar atmosphere, 4 hrs                  |
| GNF-1600C-H   | GNF-1600C sample further heat treated at 400°C, 100-sccm flow of 4at% H₂/Ar, 24hrs    |
| GNF-2000C     | GNF-1050C sample further heat treated at 2000°C, He atmosphere, 4hrs                   |
| GNF-2500C     | GNF-1050C sample further heat treated at 2500°C, He atmosphere, 4hrs                   |

indicate that the hydrogenation of graphene could convert highly conductive zero-overlap semimetal graphene into an insulator (graphane)\(^{11}\). If this occurs, hydrogen may adversely impact lithium storage due to the loss of electrical conductivity. A brief survey of the literature on graphene nanosheets made by wet chemistry and thermal exfoliation methods fails to yield any clear-cut correlations among synthetic methods, hydrogen content, and lithium storage capacities\(^{17}\), suggesting the rather complex roles of impurities and defect structures in influencing lithium storage behaviour of graphene materials\(^{18}\). Computer simulations in fact showed that perfect graphene lacks lithium storage mechanisms\(^{19}\), and that defect structures are prerequisite for lithium storage\(^{20,21}\). In practice, however, defect sites of graphene tend to bind functional groups that often contain hydrogen (e.g., hydroxyl, carboxyl, amine, hydrogen). This underscores the universal significance of understanding the roles of hydrogen in influencing the electrochemical behaviour of graphene.

To investigate the involvement of hydrogen and hydrogenated defects in the lithium storage ability of graphene, we apply various heat treatment conditions combined with hydrogen exposure and interrogate the electrochemical performance of 3D graphene nanofoam (GNF) electrodes. Our studies use self-assembled 3D graphene nanofoams due to their numerous potential applications, including hydrogen storage, catalysis, filtration, insulation, energy sorbents, capacitive desalination, supercapacitors, and LIBs\(^{22}\). The binder-free nature of graphene 3D foam makes them ideal for mechanistic studies without the complications caused by additives. We report a drastically improved rate capacity in GNFs after hydrogen treatment, which we attribute to complex interactions between defects and dissociated hydrogen that alter the chemistry and morphology of the substrate. We discuss the atomistic origins of this behaviour through a series of control experiments and multiscale simulations, and suggest its use as a strategy for optimizing lithium transport and reversible storage in graphene-based anode materials.

**Results**

**Structural and electronic characteristics.** Our electrode materials are 3D GNFs (250 μm thick, 5–10 mm in diameter, inset of Fig. 1a) constructed from graphene oxide sheets through a sol-gel process (Methods)\(^{22}\). A series of comparison GNF samples are investigated, namely, GNF-1050C, GNF-1050C-H, GNF-1050C-H\(_2\), GNF-1600C, GNF-1600C-H, GNF-2000C, and GNF-2500C. The processing conditions of each sample are shown in Table 1. Due to a large number of GNF samples involved in our studies, the data presented in this work mainly focus on three key samples that help to yield the most valuable information for computational understanding. These samples are GNF-1050C, GNF-1050C-H, and GNF-1600C. Note that samples with ‘H’ labels are those treated in H\(_2\) environment. The original 3D structure of these three GNFs can be seen in Fig. S1 of Supplementary Information (SI). The transmission electron micrograph (TEM) of the starting material GNF-1050C in Fig. 1a indicates that it consists of relatively transparent regions attributed to few-layer graphene and dark rippled areas, where occasionally stacked layers are visible. Most interlayers are observed to be twisted due to the strain that is geometrically necessary to construct 3D GNFs. The relatively rough features on the transparent graphene layers suggest the defective nature of these entities. Raman spectroscopic analysis in Fig. 1b reveals that at the laser excitation energy of 1.96 eV, GNF-1050C has a D/G band intensity ratio of 1.45 ± 0.03, consistent with the defective nature of graphene. After annealing in a hydrogen environment (4.0 at.% H\(_2\) + Ar) at 400°C (Methods), we note that the D/G band intensity ratio of graphene (GNF-1050C-H) is marginally changed (1.48 ± 0.03). However, a blue-shift of E\(_{2g}\) mode (i.e., G-band) is observed in GNF-1050C-H (inset of Fig. 1b)\(^{23}\), suggestive of additional contribution from non-zone centre phonons, likely due to disorder at domain boundaries. The energy dispersive double resonant D-band at different laser energies (SI, Fig. S2) exhibits a slope of ~60 cm\(^{-1}\)/eV for GNF-1050C and GNF-1050-H that is larger than the typical value for graphene (~38 cm\(^{-1}\)/eV)\(^{24}\). The variation in D-band frequency obtained with the same laser energy may be ascribed to a slight variation in force-constants between these two materials, with the GNF-1050C-H exhibiting tighter binding\(^{25}\).

The D/G band intensity ratio increases to 2.67 ± 0.18 when GNF is annealed at a higher temperature of 1600°C in Ar environment (GNF-1600C). The increased D/G band ratio at higher annealing temperatures is counterintuitive but suggests that the GNF-1050C and the GNF-1600C belong to Stage II
and Stage I defect region materials with an estimated graphene domain size of 1.4 nm and 10.4 nm\textsuperscript{18,26,27} (SI, Fig. S3), respectively (Methods). The Brunauer-Emmett-Teller (BET) surface area measurement and porosimetry characterization (SI, Fig. S4) reveal that H\textsubscript{2}-treatment leads to a slight shift towards smaller pores and a marginal decrease of surface area, so does the higher temperature treatment (Table 2). Elastic recoil detection analysis (ERDA) in Fig. 1c indicates that the hydrogen distribution inside GNFs is generally nonuniform with higher hydrogen on the surface and levels off after ~10\textmu m deep. The average H-content is found to increase from 3.6 at.% in GNF-1050C to 4.3 at.% in GNF-1050C-H (Methods); i.e., H\textsubscript{2} treatment increases the total atomic hydrogen level in GNFs. Interestingly, we do not observe an obvious increase of d-spacing for hydrogen treated sample (SI, Fig. S5).

The electronic structure of three GNF samples (GNF-1050C, GNF-1050-H, GNF-1060C) is investigated by carbon K-edge X-ray adsorption spectroscopy (XAS), Fig. 1d. These spectra display features characteristic of $sp^2$ carbon materials, including a sharp resonance at $\sim$285.4 eV that is attributed to C(1s) $\rightarrow$ (C-C) $\pi^*$ transitions, a feature at $\sim$291.5 eV arising from a core-hole (\sigma) exciton state\textsuperscript{28}, and a manifold of resonances at higher energies primarily associated with C(1s) $\rightarrow$ (C-C) $\sigma^*$ transitions\textsuperscript{29}. Subtle changes in the x-ray absorbance are observed following exposure of the GNF-1050C to 4.0 at.%-H\textsubscript{2} environment at 400°C. The GNF-1050C-H spectrum exhibits a slight reduction in intensity of both the C(1s) $\rightarrow$ (C-C) $\pi^*$ resonance and the C(1s) $\rightarrow$ (C-C) $\sigma^*$ resonance at $\sim$292.5 eV with respect to the GNF-1050C data. In contrast, any changes in the intensity of the C(1s) $\rightarrow$ (C-H) $\sigma^*$ resonance fall within experimental error. The diminished intensity of the C-C $\pi^*$ and $\sigma^*$ resonances suggests a reduction in the overall $sp^2$ character of the GNF\textsuperscript{29} following exposure to H\textsubscript{2}. In light of this assignment, the absence of any accompanying intensity changes in the C(1s) $\rightarrow$ (C-H) $\sigma^*$ is important because it eliminates possible mechanisms by which the proportion of $sp^2$ carbon could decrease. In particular, the addition of molecular hydrogen across two adjacent $sp^2$ hybridized carbon atoms (i.e. to yield two $sp^3$-hybridized carbons)
is omitted as a viable mechanism for the reduction in $sp^2$ character because it would necessitate the observation of a corresponding increase in intensity of the C(1s) $→$ (C-H) $\sigma^*$ resonance at $\sim$ 289.5 eV$^{29}$. For the GNF-1600C sample, we observe an increase in intensity of the C(1s) $→$ (C-C) $\pi^*$ resonance, indicative of an increase in the proportion of $sp^2$ hybridized carbon within the GNFs. This assignment is further supported by the observed sharpening and intensity increase in the core-hole exciton feature upon hydrogenation. Both features are characteristic of an increase in domain size and crystallinity of the GNF-1600C sample, consistent with Raman signatures.

**Electrochemical behaviour at various charge/discharge rates.** The specific capacities obtained at different charge/discharge rates for the three GNF samples shown in Fig. 2a indicate that H$_2$-treated sample (i.e., GNF-1050C-H) has substantially better rate capacity compared to the GNF-1050C sample, whereas the GNF-1050C-H has the worst rate performance (the voltage profiles of three samples at the 1$^{\text{st}}$ and 5$^{\text{th}}$ cycle can be seen in SI, Fig. S6). In addition, the intrinsically high capacity of GNF-1050C-H is manifested at 50 mA/g current density after 30 cycles. Since GNF-1050C-H and GNF-1050C have marginal different in terms of pore-size distribution and pore volume (Table 2), the significantly higher capacity and better rate performance in GNF-1050C-H cannot be simply attributed to pore size effects.

We define the capacity enhancement as $\delta = (C_{\text{GNF-1050C-H}} - C_{\text{GNF-1050C}})/C_{\text{GNF-1050C}} \times 100\%$, where $C_{\text{GNF-1050C-H}}$ and $C_{\text{GNF-1050C}}$ are the delithiation capacities of H$_2$-treated and reference GNF-1050C graphene samples, respectively. In Fig. 2b, we observe a $\delta$ value that varies with the charge rate and ranges from 17–43% after H$_2$ treatment. At lower rates, $\delta$ increases sharply before generally saturating at higher rates. The maximum enhancement of $\sim$43% is observed at the charge/discharge rate of 200 mA/g. This impressive improvement of rate capacity in H$_2$-treated sample is surprising, given the fact that the overall hydrogen content in GNF-1050C-H was only slightly increased (Fig. 1c), whereas the oxygen content remains identical (~2.0 at.%) (SI, Fig. S7) as determined by Rutherford backscattering spectroscopy (RBS). This strong electrochemical performance enhancement is highly reproducible and is also observed in another set of control samples that were annealed in 100% H$_2$ environment (SI, Fig. S8). The differential capacity curves of a series of GNF samples in the inset of Fig. 2b reveal two informative trends: (1) the lithium intercalation peaks (<0.5 V) shift towards lower potentials after H$_2$ treatment (i.e., compare GNF-1050C-H with GNF-1050C), suggestive of easier intercalation processes; (2) higher temperature annealing without H$_2$ leads to higher graphitization of graphene and thus stronger lithium intercalation peaks. However, the same higher temperature annealing also leads to the disappearance of multiple lithium reaction peaks in the voltage range of 1–3 V, causing low capacity in higher temperature annealed graphene materials (i.e., GNF-2000C and GNF-2500C, SI, Fig. S9). We observe that GNF-1600C has the largest initial lithiation capacity (~3182 mAh/g in Fig. 2a), but the lowest Coulombic efficiency (~28.7%), Fig. 2c. In comparison, the H$_2$-treated GNF-1050C-H graphene shows a lower initial lithiation capacity (~2677 mAh/g), but the highest first-cycle Coulombic efficiency (~39.0%). The observation of a lower Coulombic efficiency in the less-defective GNF-1600C (see Fig. S3) compared to the GNF-1050C contrasts with the popular belief that surface functional groups on graphene (especially the oxygen-containing groups) tend to cause side reactions of electrolyte and thus lower Coulombic efficiencies$^{30}$. Our RBS measurements in fact show much less oxygen content in GNF-1600C (Fig. S7). These results suggest that defective structures on graphene surface provide an overall benefit for achieving higher Coulombic efficiency despite their strong affinity to functional groups that are often cited as sources of side reactions$^{30}$. This observation is supported by the increasingly worse electrochemical performance of highly crystalline 3D graphene foams (i.e., less defective) that were heat treated at even higher temperatures of 2000–2500°C (Fig. S9).

To explore experimentally the origins of better Coulombic efficiency and high rate performance in H$_2$-treated graphene, we perform electrochemical impedance spectroscopy measurements of three samples after 35 charge/discharge cycles, Nyquist plots of which are shown in Fig. 2d. The most salient feature of the H$_2$-treated sample is a drastic reduction of charge transfer resistance vis-à-vis the other two samples, as manifested by a markedly smaller semi-circle at high frequencies for the GNF-1050C-H sample. Randles equivalent circuit modelling (Fig. S10 and Table S1) confirms this observation and suggests that double layer capacitance and electrolyte resistance of all three materials remain largely unchanged. This behaviour hints at possible roles of hydrogen in mediating charge-transfer and ion insertion processes during lithium binding/storage; e.g., by modifying the nature of interactions between graphitic layers.

| Samples    | Specific surface area (m$^2$/g) | Peak pore diameter (nm) | Pore volume (cm$^3$/g) |
|------------|---------------------------------|-------------------------|------------------------|
| GNF-1050C  | 1340                            | 6.0                     | 4.0                    |
| GNF-1050C-H| 1329                            | 4.3                     | 4.5                    |
| GNF-1600C  | 1067                            | 5.2                     | 3.9                    |

Table 2. Surface area and pore volume of three key GNF samples for electrochemical performance measurements.
Another important observation is the drastically increased solid-electrolyte interphase (SEI) resistance in GNF-1600C, due to higher degree of graphitization at higher annealing temperatures.

Capacitive contribution. More information related to lithium storage kinetics of our 3D GNFs is revealed by cyclic voltammetry (CV) data shown in Fig. 3. At a constant CV scan rate of 0.2 mV/s, Fig. 3a, we observe that the H₂-treated sample exhibits the largest current density, whereas the GNF-1050C sample has the smallest. Assuming that the current (i) follows a power law relation with the sweep rate (ν), we have

\[ i = a \nu^b \]

where a and b are materials-dependent variables. By using a ν range of 0.1–2 mV/s, we derive a b-value of 0.72 for the GNF-1050C, which increases to 0.75 for the GNF-1050C-H but decreases to 0.70 for the GNF-1600C sample. As b-values of 0.5 and 1 represent currents controlled by linear diffusion processes and surface or capacitive processes, respectively, the observation of intermediate b-values suggests both types of processes are active during lithiation. This is expected given the high surface area of the 3D GNFs used in this work. We can gain some additional physical insight by introducing a simplified model in which the two charge storage channels are functionally independent, in which case we can follow the formulation suggested in earlier studies that expresses the current (i) response at a given voltage (V) as a linear sum:

\[ i(V) = c_1 \nu + c_2 \nu^{1/2} \]

Here, \( c_2 \nu^{1/2} \) describes diffusion-controlled (battery-like) contributions and \( c_1 \nu \) encompasses remaining contributions that behave in a capacitive fashion. We emphasize that the “capacitive” contribution \( c_1 \nu \) could encompass a variety of possible processes that are not diffusion-controlled, including double layer capacitance, pseudocapacitance, and general kinetically fast charge storage at accessible sites. By determining \( c_1 \) and \( c_2 \), the individual contributions of diffusion-controlled and capacitive mechanisms within the independent model can be estimated at each sweeping rate, as illustrated in Fig. 3b. The capacitive
contribution for each material increases with sweeping rate, as one may expect. One of the most intriguing observations in Fig. 3b, however, is the enhanced capacitive contribution (and correspondingly decreased diffusion-controlled contribution) in the H₂-treated sample compared with the other two samples. For example, at the highest sweeping rate of 2 mV/s, we find that 66.4% of lithium storage comes from capacitive mechanisms in GNF-1050C-H versus 60.5% and 59% for GNF-1050C and GNF-1600C, respectively. At a fixed scan rate of 0.2 mV/s, we observe a similar trend for three materials in the voltage window of 0.25–3.2 V, Fig. 3c. Although the two-channel independent charge storage model upon which Eqn. 2 is based neglects the full complexity of actual system, it illustrates the substantially higher capacitive contribution after H₂ treatment, which is one of the main causes why GNF-1050C-H has the highest rate capacity.

Discussion

Atomistic mechanisms of high-rate capacity. The specific origin of the enhanced capacitive contribution upon H₂ treatment is not directly discernible. However, as previously discussed, no double layer capacitive enhancement is observed in the equivalent circuit models of Fig. 2d. Furthermore, the differential capacity in the inset of Fig. 2b reveals no obvious additional redox peaks that might signal new pseudocapacitive features; instead, we observe a general enhancement and shift of existing peaks towards lower voltage onsets. The results are therefore consistent with universally faster kinetics for Li diffusion and incorporation, eliminating diffusion-related bottlenecks at high rates. Experimentally, we may conclude that atomic hydrogen helps primarily to increase diffusion kinetics and reversibility rather than the density of storage sites. In other words, the capacity increases can be attributed chiefly to the increased accessibility and improved rate performance of the electrode. This is supported by the enhanced capacitive contributions in Fig. 3, as well as the fact that the capacity improvements over the untreated sample increase with current density as expected for a kinetically driven process (Fig. 2b). Moreover, the decrease in the available sites upon first cycle in Fig. 2a excludes the possibility of increased storage sites from hydrogen treatment. Rather, the kinetic improvement is almost certainly linked to a morphological change in the electrode, since the total amount of hydrogen incorporation is relatively low (Fig. S7) and yet induces a change capable of significantly affecting the behaviour of a large number of Li⁺.

Figure 3. Capacitive contributions. (a) C-V curves of three GNF samples at a scan rate of 0.2 mV/s. (b) The determination of capacitive and diffusion-controlled current contributions at certain sweep rates at 2V. (c) The capacitive contribution (grey area) in the voltage window of 0.25–3.2V at 0.2 mV/s. The percentage capacitive contributions are 28.7%, 30.9%, and 23.0% for GNF-1050C, GNF-1050C-H, and GNF-1600C, respectively.
ions simultaneously. The Nyquist plot in Fig. 2d offers valuable insight into this effect, since it indicates a decrease in the charge-transfer resistance and a simultaneously shortened diffusion pathway (demonstrated by the steep slope of the Warburg-like element at lower frequencies, indicative of finite diffusion). Importantly, the morphological change is accompanied by a chemical signature associated with a decrease in the aromatic ring structure of graphene. This is consistent with the XAS data in Fig. 1d that point to a reduction in $sp^2$ carbon character, as expected from disruptions in ring structures or loss of surface carbon. At the same time, the Raman data (Fig. 1b) indicate that these chemical changes do not appreciably affect the average domain size, implying the hydrogen-induced modification occurs at sites that are already defective (e.g., non-hexagonal rings or chemically functionalized carbon structures, likely concentrated at domain boundaries). Our control experiments provide further evidence that the effect of hydrogen incorporation into graphene is related to the presence of structural or chemical defects, as we observe little performance enhancement when further annealing GNF-1600C in a $H_2$ environment (SI, Fig. S11).

Based on the above important observations, we propose that hydrogen tends to “attack” high-energy structurally and chemically defective sites at domain boundaries. The reaction at the domain boundary follows three possible pathways: (1) cleavage of strained C-C bonds; (2) etching via successive reduction and hydrogenation of energetic carbon (e.g., to methane, ethane, ethene/ethylen, etc.); (3) reduction and hydrogenation of oxygen- or nitrogen-containing functional groups (e.g., to water or ammonia). In each case, the process is accompanied by H-termination of dangling carbon bonds. The broken bonds alleviate hydrogenation of oxygen- or nitrogen-containing functional groups (e.g., to water or ammonia) in the framework without creating additional case, the process is accompanied by H-termination of dangling carbon bonds. 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Mechanisms of dissociative hydrogenation. First, we demonstrate the possibility of dissociative hydrogenation (i.e., dissociation of molecular $H_2$ and chemisorption of atomic $H$) at defective sites on the graphene surface at 400 °C. According to a “hot hydrogen” model described in Ref. 34, as energetic $H_2$ molecules at the tail of the thermal distribution collide with the graphene surface, the proximity of their approach alleviates the need to overcome the $H_2$ high dissociation barrier (~4.12 eV)34 as long as an energetically favourable defective binding site is present. The induced polarization of $H_2$ near defective sites may also assist in low-temperature dissociation35,36. Accordingly, the necessary energetics for dissociation $H_2$ on defective binding sites need only be on the order of the bond energy difference between $H$ in $H_2$ and in defective graphene. Using relevant DFT energetics, combined with key information about our experimental conditions (Methods and SI, Fig. S12), we perform thermodynamic calculations to assess the extent of the dissociative hydrogenation process. Figure 4b shows that a 4.0 at.%-H$_2$ mixture at 400 °C will generate non-negligible H concentrations at defective binding sites, provided the binding of atomic H is sufficiently strong (see the “active” regime in the figure). Appreciable surface diffusion of atomic H can also be activated at 400 °C (see Fig. S13), suggesting the additional possibility of segregation of adsorbed surface H to high-energy domain boundaries.

Identification of candidate binding sites for dissociative hydrogenation. Importantly, the result in Fig. 4b gives an energetic lower bound for participation in dissociative hydrogenation. In particular, the defective binding sites must be exothermic for atomic $H$ by at least $\Delta E \sim 3.60 \text{eV}$. Note that in addition to energies associated with changes to chemical bonds ($E_b$), $\Delta E$ may include strain energy dissipation ($\Delta E_{strain}$) arising from the relaxation of highly curved or strained configurations at the attack sites$^{37}$, i.e., $\Delta E = E_b + \Delta E_{strain}$. The existence of such strained regions in our graphene samples is supported by direct experimental observation of rippled/twisted structures (Fig. 1a and S1). Assuming that $\Delta E_{strain}$ could account for $\sim 0.07$–0.42 eV per binding site (based on the strain energy of a rippled graphene sheet induced by 5% to 20% compressive strain$^{37}$; larger strains will further widen this window), we conclude that defective sites with $E_b > 3.53 \text{eV}$ (assuming 5% strain) or $> 3.18 \text{eV}$ (assuming 20% strain) are possible candidates for dissociative hydrogenation (Fig. 4b).
Figure 4 illustrates some of the chemical processes that could lead to the formation of σ-type C-H bonds with values of $E_B$ fitting Fig. 4b. The computed energy of H binding on a free nanoribbon edge is 4.88 eV (Fig. 4c), well within our window (by contrast, the value for basal H binding on graphene is only 0.81 eV, prohibiting dissociative hydrogenation at conventional surface sites). The target range for $E_B$ is also compatible with H binding at the edges of multi-vacancy clusters (e.g., the 5-vacancy cluster shown in Fig. 5a has an edge H binding energy of 3.55 eV). Several other possible reactions involving successive hydrogenation of domain-edge carbon atoms and of residual oxygen or nitrogen are shown in Fig. 4c; such defect motifs may be introduced during processing or upon reaction with environmental agents. Scenarios include H$_2$ dissociation to etch away atomically rough edges by formation of CH$_4$, or else the replacement of O-, OH-, or NH$_x$-containing groups with H$_2$O or NH$_3$, hydrogenating any remaining dangling bonds in the process. Note that molecular products of these reactions (e.g., H$_2$O, CH$_4$, and NH$_3$) would be released at low pressure or elevated temperatures, effectively conserving the total amount of hydrogen in the system; this may explain the largely similar H concentrations before and after hydrogen treatment detected in high vacuum conditions ($<10^{-6}$Torr) by ERDA and implied in

**Figure 4. Atomistic mechanisms of dissociative hydrogenation.** (a) Schematic of proposed mechanism for enhanced rate performance with H$_2$ treatment, in which H$_2$ dissociates at strained, functionalized, and/or highly defective domain boundaries, terminating edges and opening up the graphene sheet for improved Li penetration. (b) Equilibrium fraction of available sites terminated via a dissociative hydrogenation process of “hot” H$_2$ hydrogen saturation ($X_H$) as a function of the C-H bond formation energy at the site ($E_B$). The ranges of $E_B$ leading to active dissociative hydrogenation are highlighted for unstrained graphene (blue region), as well as with 5% strain (green dotted line and arrow) and 20% strain (purple dotted line and arrow) based on values in Ref. 37. The red dotted line indicates full saturation of binding sites considered in the model. (c) Bond formation energies $E_B$ (per H) for the dissociative reaction of H$_2$ with candidate edge functional groups with energetics sufficient for full saturation. The right side shows the corresponding hydrogenation products. The red dotted line represents hydrogenation of a reactive zigzag edge.
the XAS results. To conclude, we find that functionalized edges, multi-vacancy complexes, and strained non-hexagonal rings — likely to be concentrated at strained domain boundaries — all constitute probable sites for dissociative hydrogenation.

**Enhanced Li penetration at hydrogenated boundaries.** To test the degree of kinetic enhancement associated with hydrogenation of multi-vacancy complexes and domain boundaries according to the mechanism in Fig. 4a, we estimated the DFT barrier associated with Li diffusion through hydrogen-terminated perforations of different sizes within a pristine graphene sheet (Methods). As shown in Fig. 5a, we find that very low barriers (~0.3 eV) are obtained even for relatively small perforations (>3 atom vacancies) provided the bonds are hydrogen passivated. Accordingly, even slight opening of the lattice due to strain alleviation or carbon etching at strained domain boundaries could have significant impacts on the diffusion kinetics, with passivation mitigating any chemical or electrostatic interaction with Li\(^+\) that could otherwise lead to irreversible trapping. We point out that increases in the interlayer spacing upon insertion and binding of surface H between graphene layers could further aid intercalation near the perforations, as explored theoretically in SI, Fig. S14; however, analysis of X-ray diffraction patterns provides little evidence of systematic increases in the layer spacing upon hydrogenation of our samples (Fig. S5).

**Additional reversible capacity.** Although the dominant effect of hydrogenation is improved kinetics associated with enhanced surface capacitive behaviour (suggested by the results in Fig. 3 and the mechanism of Fig. 4a), our calculations also suggest that the addition of hydrogen at a dangling-bond or basal-plane site will have a secondary positive impact on overall capacity by activating nearby reversible binding sites (Fig. 5b and SI, Supplementary discussion, Figs S15–S19). On pristine graphene, it is well established that Li\(^+\) binding is too weak for practical use as an anode. Interestingly, the addition of hydrogen enhances the adsorption of Li\(^+\) in the hexagonal rings nearest the hydrogen (specifically, the near-edge ring for a dangling-bond edge site or a nearest-neighbour ring on the opposing side of a basal-plane site, as shown in Fig. 5b). The resulting Li binding energies are within a target range that encompasses adequate stability against Li agglomeration (\(E_b > 1.6\) eV) while maintaining reversibility.

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**Figure 5. Mechanisms of high-rate capacity enhancement.** (a) Estimated barrier for Li penetration through a graphene sheet perforated with different H-terminated pore sizes, demonstrating kinetic enhancement. (b) Binding energies (eV) of Li (green) at a pristine site (lower left), compared with binding near H atoms located at basal (top) and edge (right) sites where additional capacity can be activated.
under practical cycling conditions. The introduction of curvature would further stabilize such sites. We emphasize that these mechanisms can only add reversible capacity from local sites nearest the hydrogen, and would therefore have a negligible effect on overall kinetics. Accordingly, they cannot represent the dominant mechanism for our reported high-rate capacity increase, but may nonetheless provide a measureable contribution.

In summary, we have investigated the “hydrogen effect” in the electrochemical performance of defective and highly crystalline 3D GNFs, through a series of heat treatment experiments and computer simulations. Defects (e.g., vacancies, edges) and hydrogen are found to play an integral role in helping enhance the lithiation kinetics, leading to high rate capacity in hydrogen treated graphene materials. This behavior appears universal in all hydrogen treated samples. Among several beneficial roles of hydrogen, the binding of hydrogen along domain boundaries is considered as an effective pathway to help achieve high-rate performance 3D GNF electrodes, as it can substantially lower the barrier for Li+ penetration. Our studies further reveal a synergy between hydrogen and defects, in that hydrogenation becomes ineffective in highly crystalline GNFs due to a substantial loss of defective sites for hydrogen binding. This connection underscores the desirability of synthetic avenues or processing treatments that provide some residual native defects or O- and N-containing functional groups as target sites for hydrogenation in graphene derivatives. Questions remain as to how to optimize defect density and hydrogen incorporation in graphene materials in order to achieve high energy density and high power density for LIB applications.

Methods
Fabrication of graphene nanofoams (GNFs) and heat treatment. GNFs were synthesized by gelation of graphene oxide followed by supercritical drying and high temperature pyrolysis. 400 mg single-layer graphene oxide powder (~300–800 nm in lateral dimensions, Cheap Tubes inc.) were suspended in 20 mL deionized (DI) water (resistivity of 18.2 MΩ). After adding 4.22 ml concentrated ammonium, the suspension was sonicated overnight at 15 °C. The obtained slurry was poured into a rubber mould in 20 mL deionized (DI) water (resistivity of 18.2 MΩ). The wet gel was washed in DI water, followed by acetone. The acetone was exchanged with liquid CO2 in a critical point drier overnight. The drier was heated to 50 °C and the pressure was adjusted to maintain ~1500 psi to exceed the critical point of CO2. The gel was held at this state for 2h, after which the drier was slowly vented to ambient pressure to remove dried gels. The obtained dried gels were placed in a tube furnace for carbonization at 1050 °C for 4h in N2. The GNF sample was further annealed at 400 °C in 4 at% H2/Argon for 24h to obtain GNF-1050C-H, or pure H2 for 4h to obtain GNF-1050C-H2. To increase crystallinity, GNF-1050C was further annealed at 1600 °C in argon, 2000 °C in helium, and 2500 °C in helium for 4h.

SEM and TEM characterizations. Scanning electron microscopy (SEM) was conducted using a JOEL JSM-7401F field-emission SEM under secondary electron imaging (SEI) mode with an accelerating voltage of 20 kV and a beam current of 20 mA. The cross sectional GNFs with fresh fracture surface was examined. Transmission electron microscopy (TEM) was performed on a Philips CM30 field-emission TEM. Samples were prepared by manually crashing GNF samples onto a Lacey carbon supported copper grids (200 mesh, Ted Pella, Inc.).

Porosimetry. The porosimetry was determined by Brunauer-Emmett-Teller (BET) and Barrett-Joyner-Halenda (BJH) methods in nitrogen adsorption/desorption experiments using an ASAP 2020 Surface Area Analyser (Micromeritics Instrument Corporation). GNF samples with mass of 10 mg to 20 mg were preheated at 300 °C under N2 flow for 24h to remove moisture.

Raman spectroscopy. Raman spectra were collected using three laser excitation wavelengths, 473 nm, 532 nm, and 633 nm. The spot size was 1.0–2.5 μm, and the laser beam power was 1–2 mW. Table S2 lists the detailed spot size, power, and spectral resolution of each excitation laser wavelength for the measurements. At least three spectra were collected for each sample. To analyse the position and intensity of D, G, and D’ bands, the background or raw spectra data was subtracted by a linear function and the peaks were deconvoluted by multi-peak fitting using Lorentzian functions. The domain size or crystallite size (L_D) was estimated from I(D)/I(G) ratio using the following equation:

\[
\frac{I(D)}{I(G)} = C_A \frac{(r_A^2 - r_E^2)}{(r_A^2 - 2r_E^2)} \cdot \frac{\pi r_E^2}{L_D^2} - e^{-\frac{\pi (r_A^2 - r_E^2)}{L_D^2}}
\]

Here \(C_A\) is the weighing parameter of an active area (A) that can be expressed as \(C_A = AE_{el}^{-1}\), where \(A = (160 ± 48)\) eV, B = 4, and \(E_{el} = 1.96 eV\). \(r_A\) and \(r_E\) are two length scale parameters that determine the region of D band scattering, which were fitted to be 3 nm and 1 nm, respectively.
X-ray absorption spectroscopy (XAS). XAS measurements were performed at beamline (BL) 8.0.1.1 of the Advanced Light Source, Lawrence Berkeley National Laboratory. Carbon K-edge XAS data was recorded in the total electron yield (TEY) mode over a spectral range of 280–330 eV, with the sample angled at 45° with respect to the incident x-ray beam. Energy calibration of BL8.0.1.1 was achieved via reference to the C(1s) → π* resonance for a freshly cleaved sample of highly oriented pyrolytic graphite (285.38 eV)49. All XAS data was normalized to both the incident x-ray flux, I0, and the absorption edge step, I0 was measured concurrently with the XAS signal via the drainage current from a gold grid located upstream of the experimental sample and the absorption edge step was taken as the difference in absorbance in the pre-edge (280 eV) and post-edge (330 eV) regions of the XAS spectrum.

X-ray diffraction (XRD). XRD was carried out using Bruker AXS D8 ADVANCE X-ray diffractometer with Cu radiation source operated at 40 kV and 40 mA. 2θ scan was conducted from 10° to 90° with 0.02° steps and 2 s counting time per step.

Elastic recoil detection analysis (ERDA) and Rutherford back scattering (RBS). Sample was mounted on a Si piece. Mounting and sample transfer were completed in N2 environment. The H-content discussed in Ref. 34, the dissociation reaction can be written as

\[ \Delta G = -\beta \cdot \left[ C - H(s) \right] \rightarrow \left[ C - H(s) \right] + \beta \cdot \left[ H_2(g) \right], \]

where \( \beta \) is the extent of reaction and \( H_2(g) \) represents a ‘hot’ hydrogen gas molecule. The moles of \( H_2 \) at temperature \( T \) is calculated by

\[ n_{H_2}^0 = n_{He}^0 \exp(-E_D/k_B T), \]

where \( n_{He}^0 \) is the total moles of \( H_2 \) molecules flowing into the chamber and \( E_D = [\Delta E - |E_b|] \). The total energy variation upon dissociative hydrogenation is computed as follows:

\[ \Delta G = \beta \cdot \left[ \mu_{H}^0 - (1/2) \mu_{H_2}^0 \right] + \Delta G_{H_2}^{\text{pressure}} + G_{C/H}^{\text{mix}} + \beta \cdot N_0 \cdot \Delta E, \]

where \( \mu_{H}^0 \) and \( \mu_{H_2}^0 \) are the molar Gibbs free energies of H atoms and \( H_2 \) molecules, respectively, at the standard state, and \( N_0 \) is Avogadro’s number. \( \Delta G_{H_2}^{\text{pressure}} \) is the free energy change upon reaction due to the partial pressure variation (accounting for zero-point, entropy, and enthalpy contributions) and \( G_{C/H}^{\text{mix}} \) is the free energy of mixing for the hydrogenated graphene. Ideal-gas and ideal-mixing behaviour are assumed. Using Eq. (5) and the phase fraction definitions in Table S3, we determine the \( \beta \) that minimizes \( \Delta G \) to obtain the thermodynamically stable H content. See Supplementary Methods for further details.

First-principles simulations. We use density functional theory implemented in VASP46 with the projector augmented wave method47 and the Perdew-Burke-Erzenhof (PBE) exchange-correlation functional48. Basal-plane Li adsorption was simulated using a 6 × 6 graphene supercell. Edge-site Li adsorption was simulated using a zigzag graphene nanoribbon in a 72-atom orthorhombic unit cell, edge-terminated with 12 hydrogen atoms. A 400 eV plane-wave cutoff was used. Brillouin zone sampling for the basal (nanoribbon) case was based on a 5 × 5 × 1 (5 × 1 × 1) Monkhorst-Pack k-point mesh. Periodic boundary conditions were used, with a 20 Å vacuum layer inserted perpendicular to the planes and a dipole correction applied. Gaussian smearing was used with a smearing width
of 0.03 eV. Forces were converged to $<1 \times 10^{-2}$ eV/Å. For calculation of surface diffusion barriers (Fig. S11), the nudged elastic band (NEB) method was used. Li binding energies were calculated as

$$E_{\text{bind}} = [E_{\text{tot}}(X) + E_{\text{tot}}(\text{Li}) - E_{\text{tot}}(X + \text{Li})] / N_{\text{Li}},$$

where Li, X, and X + Li refer to an isolated Li atom and to Li-free and Li-adsorbed substrates, respectively, and $N_{\text{Li}}$ is the number of Li atoms in the cell. Vacancy complexes were generated by terminating dangling bonds with hydrogen within the graphene plane. The energy barrier for Li penetration through the vacancy complexes was estimated as the energy differences between constrained optimizations of two Li positions: (1) above the largest opening in the graphene sheet at a distance of 1.72 Å (the equilibrium binding distance of Li on the basal plane of pristine graphene); and (2) in the plane of graphene at the centre of the largest opening.

References

1. Raccichini, R., Varzi, A., Passerini, S. & Scrosati, B. The role of graphene for electrochemical energy storage. Nat. Mater. 14, 271–279, doi: 10.1038/NMAT4170 (2015).
2. Bonaccorso, F. et al. Graphene: related two dimensional crystals, and hybrid systems for energy conversion and storage. Science 347, 1246501, doi: 10.1126/science.1246501 (2015).
3. Xu, Y. et al. Holey graphene frameworks for highly efficient capacitive energy storage. Nat. Commun. 5, 4554, doi: 10.1038/ncomms5554 (2014).
4. Yang, X., Cheng, C., Wang, Y., Qiu, L. & Li, D. Liquid-Mediated Dense Integration of Graphene Materials for Compact Capacitive Energy Storage. Science 341, 534–537, doi: 10.1126/science.1239089 (2013).
5. Zhu, Y. et al. Carbon-Based Supercapacitors Produced by Activation of Graphene. Science 332, 1537–1541, doi: 10.1126/science.1200770 (2011).
6. Li, N., Chen, Z., Ren, W., Li, F. & Cheng, H.-M. Flexible graphene-based lithium ion batteries with ultrafast charge and discharge rates. Proc. Natl. Acad. Sci. USA 109, 17360–17365, doi: 10.1073/pnas.1204827109 (2012).
7. Xu, Y. et al. Solvated graphene frameworks as high-performance anodes for lithium-ion batteries. Angew. Chem. Int. Ed. 54, 1–7 (2015).
8. Park, S. & Ruoff, R. S. Chemical methods for the production of graphenes. Nat. Nanotech. 4, 217–224, doi: 10.1038/nnano.2009.58 (2009).
9. Hernandez, Y. et al. High yield production of graphene by liquid-phase exfoliation of graphite. Nat. Nanotech. 3, 563–568, doi: 10.1038/nnano.2008.215 (2008).
10. Li, Z. et al. Effect of airborne contaminants on the wettability of supported graphene and graphite. Nat. Mater. 12, 925–931, doi: 10.1038/nmat3709 (2013).
11. El-Aasser, M. S. et al. Control of graphene’s Properties by Reversible Hydrogenation: Evidence for Graphane. Science 323, 610–613, doi: 10.1126/science.1167130 (2009).
12. Ryu, S. et al. Reversible Basal Plane Hydrogenation of Graphene. Nano Lett. 8, 4597–4602, doi: 10.1021/nl802940s (2008).
13. Balog, R. et al. Bandgap opening in graphene induced by patterned hydrogen adsorption. Nat. Mater. 9, 315–319, doi: 10.1038/nmat2710 (2010).
14. Dahn, J. R., Zheng, T., Liu, Y. H. & Xue, J. S. Mechanisms for lithium insertion in carbonaceous materials. Science 270, 590–593, doi: 10.1126/science.270.5236.590 (1995).
15. Cabaña, I., López, M. I. & Alonso, J. A. Enhancement of hydrogen physisorption on graphene and carbon nanotubes by Li doping. J. Chem. Phys. 123, doi: 10.1021/jp3121572 (2005).
16. Dimitrakakis, G. K., Tylianakis, E. & Froudakis, G. E. Pillared Graphene: A New 3-D Network Nanostructure for Enhanced Hydrogen Storage. Nano Lett. 8, 3166–3170, doi: 10.1021/nl081417w (2008).
17. Vargas, O. A., Caballero, A. & Morales, J. Can the performance of graphene nanosheets for lithium storage in Li-ion batteries be predicted? Nanoscale 4, 2083–2092, doi: 10.1039/c2nr11936f (2012).
18. Ye, J. C. et al. Enhanced electrochemical performance of ion-beam-treated 3D graphene aerogels for lithium ion batteries. Carbon 85, 269–276, doi: 10.1016/j.carbon.2014.12.097 (2015).
19. Lee, E. & Persson, K. A. Li Absorption and Intercalation in Single Layer Graphene and Few Layer Graphene by First Principles. Nano Lett. 12, 4624–4628, doi: 10.1021/nl3019164 (2012).
20. Liu, Y., Artyukhov, V. I., Liu, M., Harutyunyan, A. R. & Yakobson, B. I. Feasibility of Lithium Storage on Graphene and Its Derivatives. J. Phys. Chem. Lett. 4, 1737–1742, doi: 10.1021/jz400491b (2013).
21. Liu, Y., Wang, Y. M., Yakobson, B. I. & Wood, B. C. Assessing Carbon-Based Anodes for Lithium-Ion Batteries: A Universal Description of Charge-Transfer Binding. Phys. Rev. Lett. 113, 028304, doi: 10.1103/PhysRevLett.113.028304 (2014).
22. Worsley, M. A. et al. Synthesis and characterization of highly crystalline graphene aerogels. ACS Nano 8, 11013–11022, doi: 0.1021/nn305335u (2014).
23. Ferrari, A. C. et al. Raman spectrum of graphene and graphene layers. Phys. Rev. Lett. 97, 187401, doi: 10.1103/PhysRevLett.97.187401 (2006).
24. Narula, R. & Reich, S. Double resonant Raman spectra in graphene and graphite: A two-dimensional explanation of the Raman amplitude. Phys. Rev. B 78, 165422, doi: 10.1103/PhysRevB.78.165422 (2008).
25. Matthews, M. I., Pimenta, M. A., Dresselhaus, G., Dresselhaus, M. S. & Endo, M. Origin of dispersive effects of the Raman D band in carbon materials. Phys. Rev. B 59, R6585–R6588, doi: 10.1103/PhysRevB.59.R6585 (1999).
26. Eckmann, A. et al. Probing the Nature of Defects in Graphene by Raman Spectroscopy. Nano Lett. 12, 3925–3930, doi: 10.1021/nl300901a (2012).
27. Zandiastashbar, A. et al. Effect of defects on the intrinsic strength and stiffness of graphene. Nat. Commun. 5, 3186, doi: 10.1038/ncomms4186 (2014).
28. Ma, Y. et al. Core excitons and vibronic coupling in diamond and graphite. Phys. Rev. Lett. 71, 3725–3728, doi: 10.1103/PhysRevLett.71.3725 (1993).
29. Coffman, F. L. et al. Near-edge x-ray absorption of carbon materials for determining bond hybridization In mixed sp2/sp3 bonded materials. Appl. Phys. Lett. 69, 568–570, doi: 10.1063/1.1177896 (1996).
30. Wu, Z.-S., Ren, W., Xu, L., Li, F. & Cheng, H.-M. Doped Graphene Sheets As Anode Materials with Superhigh Rate and Large Capacity for Lithium Ion Batteries. ACS Nano 5, 5465–5471, doi: 10.1021/nn0906249 (2011).
31. Goodenough, J. B. & Kim, Y. Challenges for Rechargeable Li Batteries. Chem. Mater. 22, 587–603, doi: 10.1021/cm100452z (2010).
32. Augustyn, V. et al. High-rate electrochemical energy storage through Li⁺ intercalation pseudocapacitance. Nat. Mater. 12, 518–522, doi: 10.1038/nmat3601 (2013).
33. Augustyn, V., Simon, P. & Dunn, B. Pseudocapacitive oxide materials for high-rate electrochemical energy storage. Energy Environ. Sci. 7, 1597–1614, doi: 10.1039/c3ee4164d (2014).
34. Mikoushkin, V. M. et al. Graphene hydrogenation by molecular hydrogen in the process of graphene oxide thermal reduction. Appl. Phys. Lett. 102, 4, doi: 10.1063/1.473584 (2013).
35. Kim, B. H. et al. N-type graphene induced by dissociative H-2 adsorption at room temperature. Sci. Rep. 2, doi: 10.1038/srep00690 (2012).
36. Ulman, K., Bhamuk, D., Wood, B. C. & Narasiminahan, S. Physical origins of weak H-2 binding on carbon nanomaterials: Insight from ab initio studies of chemically functionalized graphene nanoplatelets. J. Phys. Chem. C 118, 7741–7750, doi: 10.1021/jp411338a (2014).
37. Dutta, D., Wood, B. C., Bhide, S. Y., Ayappa, K. G. & Narasiminhan, S. Enhanced Gas Adsorption on Graphitic Substrates via Defects and Local Curvature: A Density Functional Theory Study. J. Phys. Chem. C 118, 7741–7750, doi: 10.1021/jp411338a (2014).
38. Acik, M. et al. The Role of Intercalated Water in Multilayered Graphene Oxide. ACS Nano 4, 5861–5868, doi: 10.1021/nn101844t (2010).
39. Batson, P. E. Carbon-1s near-edge-adsorption find-structure in graphite. Phys. Rev. B 48, 2608–2610, doi: 10.1103/PhysRevB.48.2608 (1993).
40. Tirira, J., Serruys, Y. & Trocctier, P. Forward Recoil Spectrometry. (Springer US, 1996).
41. Doolittle, L. R. Nuclear Instruments and Methods in Physics Research Section B 9, 344 (1985).
42. Baglin, J. E. E., Kellock, A. J., Crockett, M. A. & Shih, A. H. Nuclear Instruments and Methods in Physics Research Section B 64, 469 (1992).
43. Bhardwaj, T., Antic, A., Pavan, B., Barone, V. & Fahlman, B. D. Enhanced Electrochemical Lithium Storage by Graphene Nanoribbons. J. Am. Chem. Soc. 132, 12356–12358, doi: 10.1021/ja1061622 (2010).
44. Yoo, E. et al. Large Reversible Li Storage of Graphene Nanosheet Families for Use in Rechargeable Lithium Ion Batteries. Nano Lett. 8, 2277–2282, doi: 10.1021/nl800957b (2008).
45. Chase, M. W., Jr. NIST-JANAF Thermochemical Tables. J. Phys. Chem. Ref. Data Monograph 9, 1–1951 (1998).
46. Kresse, G. & Furthmuller, J. Efficient iterative schemes for ab initio total-energy calculations using a plane-wave basis set. Phys. Rev. B 54, 11169–11186, doi: 10.1103/PhysRevB.54.11169 (1996).
47. Blochl, P. E. Projector augmented-wave method. Phys. Rev. B 50, 17953–17979, doi: 10.1103/PhysRevB.50.17953 (1994).
48. Perdew, J. P., Burke, K. & Ernzerhof, M. Generalized gradient approximation made simple. Phys. Rev. Lett. 77, 3865–3868, doi: 10.1103/PhysRevLett.77.3865 (1996).
49. Henkelman, G., Uberuaga, B. P. & Jonsson, H. A climbing image nudged elastic band method for finding saddle points and minimum energy paths. J. Chem. Phys. 113, 9901–9904, doi: 10.1063/1.1329672 (2000).

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Author Contributions
J.C.Y. and Y.M.W. conceived and designed experiments. M.T.O., T.W.H., Y.Y.L. and B.C.W. initiated and performed simulations. P.C. and M.A.W. synthesized all GNF samples. J.C.Y. and Y.M.W. prepared an initial draft of the manuscript with input from all co-authors.

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