Graphdiyne as a high-capacity lithium ion battery anode material

Byungryul Jang,¹ Jahyun Koo,¹ Minwoo Park,¹ Hosik Lee,² Jaewook Nam,³ Yongkyung Kwon,¹ and Hoonkyung Lee¹,a)
¹School of Physics, Konkuk University, Seoul 143-701, South Korea
²School of Mechanical and Advanced Materials Engineering, Ulsan National Institute of Science and Technology (UNIST), Ulsan 689-798, South Korea
³School of Chemical Engineering, Sungkyunkwan University, Suwon 300, South Korea

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Using the first-principles calculations, we explored the feasibility of using graphdiyne, a 2D layer of sp and sp² hybrid carbon networks, as lithium ion battery anodes. We found that the composite of the Li-intercalated multilayer x-graphdiyne was C₆Li7.31 and that the calculated voltage was suitable for the anode. The practical specific/volumetric capacities can reach up to 2719 mAh g⁻¹/2032 mAh cm⁻³, much greater than the values of ~372 mAh g⁻¹/818 mAh cm⁻³, ~1117 mAh g⁻¹/1589 mAh cm⁻³, and ~744 mAh g⁻¹ for graphite, graphynes, and γ-graphdiyne, respectively. Our calculations suggest that multilayer x-graphdiyne can serve as a promising high-capacity lithium ion battery anode.

Lithium ion batteries are currently used in various portable electronics and are expected to be used in electric vehicles because it has a higher energy density than other rechargeable batteries. In these cases, bulk lithium metal was used as the anode. However, lithium metal anodes have some stability issues such as dendrite formation.¹,² Graphite has been shown to have high stability as an anode material because of its layered structure leading to high diffusivity of lithium ions and high stability through lithium intercalation. The maximum composite of Li-intercalated graphite is known to be C₆Li,⁴–⁵ corresponding to the specific and the volumetric capacities as large as of 372 mAh g⁻¹ and 818 mAh cm⁻³, respectively. However, the search for new anode materials that have higher capacity has continued. The following minimum requirements must be met in order to qualify as an anode material: (i) the open circuit voltage (OCV) of the battery should be in the range of ~0–2 V when bulk Li is used as a reference electrode,² (ii) the specific (volumetric) capacity should be larger than that of graphite, 372 mAh g⁻¹ (818 mAh cm⁻³),² (iii) the volume expansion from lithium insertion should be as small as that for graphite. However, segregation of lithium takes place if the voltage is negative, and a candidate anode material will no longer work (though such a material may work as a cathode) if the voltage is greater than ~2 V.

Recently, it has been reported that graphynes, 2D atomic layers consisting of two adjacent sp-bonded carbon atoms and sp²-bonded carbon atoms, exhibit intriguing electronic properties such as symmetric as well as asymmetric Dirac cones.⁶–¹⁰ Another attractive feature is that the surface area of graphene is larger than that of graphene because its hexagonal area is much larger. From an application point of view, the structural properties of graphene could allow a variety of potential applications for energy storage. For instance, it has been found that carbon-decorated carbon chain networks and calcium/lithium-decorated graphynes can be used as high-capacity hydrogen storage material due to the large surface area.¹¹–¹⁴ Higher Li gravimetric density than the sp²-bonded carbon network is possible due to a larger number of Li adsorption sites. In fact, recent theoretical studies¹⁵,¹⁶ showed significant increases in both specific and volumetric capacity of Li-intercalated multilayer x- and γ-graphynes have been found; a specific capacity of 1117 mAh g⁻¹ (three times that of graphite) and the volumetric capacity of 1364 and 1589 mAh cm⁻³ (cf. ~818 mAh cm⁻³ for graphite) have been suggested.¹⁶ While a bottom-up approach¹⁷,¹⁸ was proposed to synthesize sp-sp² carbon networks such as graphynes, graphdiyne, which is a 2D atomic layer consisting of four adjacent sp-bonded carbon atoms and sp²-bonded carbon atoms, was synthesized in the form of films.¹⁹ Some experiments show that new carbon allotropes of sp-sp² hybrid carbon networks could be synthesized in the immediate future. Since graphdiyne has more sp-bonded-carbon atoms than graphyne, graphdiyne has larger surface area than graphyne. Therefore, higher Li capacity is expected for graphdiyne than for graphyne.

Recently, Sun et al.²⁰ explored possible applications of a single-layer γ-graphdiyne for lithium storage by using the density-functional calculations. They found high Li mobility in multilayer γ-graphdiyne and Li intercalation density as high as C₆Li₂. x-graphdiyne is a different graphdiyne with a honeycomb lattice like graphene, so its surface area for intercalation of lithium ions may be the largest because its surface area larger than that of other types of graphdiynes. On the other hand, since x-graphdiyne is metallic while γ-graphdiyne is semiconducting with the density functional theory (DFT) bandgap of 0.53 eV,²¹,²² x-graphdiyne would be advantageous in terms of the electronic conductivity requirement for anodes. In this study, we explored the feasibility of monolayer or multilayer x-graphdiyne for lithium ion battery anodes because the different symmetries and local geometries could lead to different lithium intercalation.

We found that the voltage is suitable for the anodes and

¹Author to whom correspondence should be addressed. Electronic mail: hkiee3@konkuk.ac.kr
the maximum composite of Li-intercalated multilayer \( \chi \)-graphdiynes is \( \text{C}_6\text{Li}_{7.31} \). This corresponds to specific/volumetric capacities of 2719 mAh g\(^{-1}\)/2032 mAh cm\(^{-3}\), much greater than the values of ~372 mAh g\(^{-1}\)/~818 mAh cm\(^{-3}\), ~1117 mAh g\(^{-1}\)/~1589 mAh cm\(^{-3}\), and ~744 mAh g\(^{-1}\) of graphite,\(^1\) graphynes,\(^{16}\) and \( \chi \)-graphdiyne,\(^{20}\) respectively. Our calculations suggest that multilayer \( \chi \)-graphdiyne can serve as high-capacity lithium ion battery anodes.

Our calculations were performed using a first-principles method based on density functional theory\(^{23}\) as implemented in the Vienna Ab-initio Simulation Package (VASP) with a projector-augmented-wave (PAW) method.\(^{24}\) The exchange correlation energy functional was used with the generalized gradient approximation (GGA) in the Perdew–Burke–Ernzerhof scheme,\(^{25}\) and the kinetic energy cutoff was set at 400 eV. Our model \( \chi \)-graphdiyne system was a \( 1 \times 1 \) hexagonal cell containing 14 C atoms. A geometrical optimization of Li-adsorbed \( \chi \)-graphyne was carried out within a fixed \( 1 \times 1 \) cell obtained from the equilibrium lattice constant of the isolated graphdiyne until the Hellmann–Feynman force acting on each atom was less than 0.01 eV/Å. The first Brillouin zone integration was done with the GGA-D2 method,\(^{27}\) which describes the interlayer van der Waals interaction including the nearest C atom is \( \sim 1.88 \text{ eV/Li.} \)

At higher concentration, the favorable site is slightly off center of the edge, as shown in Figure 1(a) when each hexagon accommodates one Li atom. Unlike multilayer \( \chi \)-graphdiyne that Li atoms are attached with the in-plane configuration,\(^{20}\) when one Li atom is added further, out-of-plane adsorption of Li atoms gets in to favorable energy as shown in Figure 1(b). The binding energy of Li atoms on a \( 1 \times 1 \) \( \chi \)-graphdiyne was calculated as a function of the Li concentration \( x \), where \( x \) is defined in \( \text{C}_6\text{Li}_x \). Here, the binding energy of Li atoms is defined by \( E_{\text{bind}}^x (\text{Li}) = (E_C + N \cdot E_{\text{Li}} - E_{\text{C-Li}}^x)/N \), where \( N \) is the number of Li atoms attached per \( 1 \times 1 \) cell for a given \( x \). \( E_{\text{C-Li}}^x \) is the total energy of Li-dispersed in a \( 1 \times 1 \) graphdiyne with a \( x \) concentration of Li atoms, \( E_C \) is the total energy of an \( 1 \times 1 \) isolated \( \chi \)-graphdiyne, and \( E_{\text{Li}} \) is the total energy of an isolated Li atom in a vacuum.

The composites of Li-dispersed \( \chi \)-graphdiyne shown in Figure 1 are (a) \( \text{C}_6\text{Li}_{1.43} \) and (b) \( \text{C}_6\text{Li}_{1.86} \), respectively. Three adsorption geometries at the composite of \( \text{C}_6\text{Li}_{1.43} \) are shown in Figure 1(a) when each hexagon accommodates one Li atom. The most favorable adsorption site is out-of-plane and slightly off center of the edge of the hexagon, as shown in Figure 1(a), where the distance between the Li atom and the nearest C atom is \( \sim 2.12 \text{ Å and the Li binding energy is } \sim 1.88 \text{ eV/Li.} \) At higher concentration, the favorable site is out-of-plane and slightly off center of the edge as shown in Figure 1(b) when each hexagon accommodates two Li atoms. This is consistent with lower concentration case. The Li binding energies on \( \chi \)-graphdiyne are greater than the cohesive energy (\( \sim 1.6 \text{ eV} \)) of bulk Li.\(^{16}\) From this, we expect

**FIG. 1.** Cross section view and side view for three different atomic structures of Li atoms attached on single layer \( \chi \)-graphdiyne at the (a) \( x = 0.43 \) and (b) \( x = 0.86 \) concentrations. The grey and purple dots represent carbon atoms and lithium atoms, respectively. The total energy \( E \) for the geometries is presented, where the energy for the lowest configuration is set to zero.
that Li atoms can be dispersed on $\pi$-graphdiyne without segregation of Li. The calculated binding energies of Li atoms on the graphdiyne are smaller than the corresponding GGA values of 2.21 eV/Li (2.69 eV/Li) on $\pi$-(\gamma)graphyne, but larger than 1.10 eV/Li on graphene, 1.80 eV/Li on a C$_{60}$, and 0.34 (0.41) eV/Li on the inner (outer) shell of a (5,5) carbon nanotube. 30 The value is comparable to 1.83 eV/Li on carbyne. 31 The binding energy of Li atoms may be increased by enhancing the electrostatic interaction between the Li atom and the C- atom due to their reduced dimensionality compared to other materials.

We carried out calculations on the adsorption of Li atoms on the graphdiyne as the concentration $x$ increases. When four Li atoms are placed on each hexagon, two Li atoms are attached with an in-plane configuration and the other Li atoms undergo out-of-plane adsorption (Figure 2(a)). In general, out-of-plane adsorption of Li atoms is preferred as the concentration becomes higher. We also found that the sp-bonded carbon atoms are distorted by Li adsorption like a zigzag carbon chain and each sp-bonded carbon atom adsorbs two Li atoms as shown in Figure 2(d). This is similar to the geometry of Li adsorption to carbyne. 31

![Image](image.png)

**FIG. 2.** The cross section view and side view for the optimized atomic structures of Li attached to single-layer $\pi$-graphdiyne at the different concentrations, (a) C$_6$Li$_{1.72}$, (b) C$_6$Li$_{3.87}$, (c) C$_6$Li$_{6.60}$, and (d) C$_6$Li$_{11.18}$. (e) The calculated average voltage per Li atom as a function of the concentration $x$.

but is in contrast to Li adsorption on $\pi$-graphyne where Li atoms are attached to in-plane hexagons with some distortion along the in-plane direction. To investigate the usability of Li-dispersed $\pi$-graphdiyne for anodes, the open circuit voltage ($V_{OCV}$) is calculated as a function of $x$

$$V_{OCV}(x) = \frac{E_C + N \cdot E_{Li}^{bulk} - E_{C-Li}^x}{eN},$$

where $e$ is the charge of electron and $E_{Li}^{bulk}$ is the total energy per Li of bcc lithium. As shown in Figure 2(e), $V_{OCV}$ decreases from $\sim$0.5 V to $\sim$0.1 V as the concentration increases as shown in Figure 2(e), which is due to the stronger Li-Li repulsive interaction at higher concentrations. The calculated $V_{OCV}$ is in the window of the required voltage for anodes. Since the added voltage is negative when the concentration is over $\sim$11.20, we set the maximum composite of the Li-dispersed graphdiyne to be C$_6$Li$_{11.18}$, corresponding to the specific capacity of 4259 mAh g$^{-1}$. This is comparable to the value $\sim$4800 mAh g$^{-1}$ of Li-dispersed carbyne. 33 This result shows that as far as the Li adsorption is concerned, $\pi$-graphdiyne acts as a two-dimensional carbon chain network.

A layered structure such as graphite is desirable for anode materials because of the diffusivity of Li, the small volume change through Li intercalation between the layers, long life cycle by good reversibility of Li insertion and extraction, and good scalability. So, multilayer graphdiynes could be a possible replacement for lithium ion battery anodes. Since AB stacking multilayer $\pi$-graphdiyne is energetically favorable, we performed calculations on Li intercalation to AB stacking multilayer $\pi$-graphdiyne. From the calculations of Li dispersion on single-layer $\pi$-graphdiyne discussed above, we predicted stable Li-intercalated multilayered $\pi$-graphdiyne structures. Figure 3 shows the optimized geometries of Li-intercalated multilayer $\pi$-graphdiyne along with the voltage of Li atoms for different concentrations of $x$ with different interlayer distance from 2.75 Å to 3.75 Å. The lowest energy interlayer distance was found to be 3.25 Å near high Li concentrations as shown in Figure 3(a). Figure 4 shows the cross section and side views, respectively, of Li-intercalated AB stacking $\pi$-graphdiyne with different concentrations where the layer-to-layer distance is 3.25 Å. The Li atoms are intercalated between the layers, and the local geometry of the adsorbed Li atoms is similar to that in Li-dispersed carbyne. 31 The computed equilibrium layer-to-layer distance of the AB-stacked $\pi$-graphdiyne estimated with the GGA-D2 calculation is 3.15 Å. Therefore, as a result of the Li intercalation, the interlayer distance increases from 3.15 Å to 3.25 Å, while the lattice constant along with the in-plane direction is unchanged. This corresponds to the volume expansion of $\sim$3%, which is smaller than that of $\sim$12% of Li-intercalated graphite where the interlayer distance increases from 3.35 Å to 3.76 Å. 32 So, the requirement of the volume change by Li intercalation is met.

Since the OCV on multilayer $\pi$-graphdiyne is positive in the concentration range as shown in Figure 3(b), it is concluded that Li atoms are intercalated without segregation within the range. However, we chose $x = 7.31$ for the maximum Li concentration because the further intercalation process is endothermic. Therefore, the maximum configuration...
expected for Li-intercalated multilayer $x$-graphdiyne is $C_6Li_{7.31}$. This composite corresponds to the practical specific capacity of $2719 \text{mAh g}^{-1}$, which is significantly greater than $372 \text{mAh g}^{-1}$ and $1117 \text{mAh g}^{-1}$ of Li-intercalated graphite and multilayer graphynes with the composite of $C_6Li_3$, respectively. In addition, the calculated practical volumetric capacity of Li-intercalated multilayer $x$-graphdiyne ($C_6Li_{7.31}$) is $2032 \text{mAh cm}^{-3}$, which is much greater than the capacity of graphite ($\sim 818 \text{mAh cm}^{-3}$) and multilayer graphynes ($1364 \text{mAh cm}^{-3}$ and $1589 \text{mAh cm}^{-3}$ for $x$- and $\gamma$-graphynes with the composite of $C_6Li_3$). The results suggest that multilayer $x$-graphdiyne can serve as high-capacity anode materials.

We have investigated energetics and geometry optimization of Li-dispersed monolayer and Li-intercalated AB-stacked multilayer $x$-graphdiyne as a promising anode material. The sp-bonded carbon atoms of $x$-graphdiyne bind Li atoms resulting in a desirable voltage for anodes and the local adsorption geometry, i.e., a zigzag chain, is similar to that of a one-dimensional carbon chain of carbyne. Four-adjacent-sp-bonded carbon chain is the minimum length mimicking carbyne. Since Li adsorption geometries are similar to those of carbyne, the Li capacity of multilayer $x$-graphdiyne may be highest among 2D layers of sp-sp$^2$ hybrid carbon networks or when compared to other carbon nanostructures consisting of sp$^2$-bonded carbon atoms. This means that $x$-graphdiyne acts like a 2D carbon chain. The volumetric capacity of 2D sp-sp$^2$ hybrid networks is significantly reduced as the length of the sp-bonded carbon chains increases. Thus, $x$-graphdiyne is an optimal candidate for lithium ion anodes when both specific and volumetric capacities are considered. We note that the structural stability of $x$-graphdiyne in the process of Li intercalation/deintercalation is critical for their practical implementations as anodes, especially considering that the calculated cohesive energies of $x$- and $\gamma$-graphdiynes (7.0 eV and 7.2 eV, respectively) or $\gamma$-graphene (7.3 eV), are smaller than that of graphite ($\sim 8.0 \text{eV}$). These values show the possibility that phase change from graphdiyne to graphite might occur during Li intercalation and deintercalation. Thus, an independent study on the stability during the electrochemical reaction will be necessary to verify them for the anode applications.

In conclusion, we performed total energy electronic structure calculations on Li-intercalated multilayer graphdiynes, using first-principles density-functional theory, to explore their applicability as lithium ion battery anodes. The calculated voltage was found to be in a desirable voltage range of the anodes. The practical specific and volumetric Li capacity of multilayer $x$-graphdiyne can reach up to $\sim 2700 \text{mAh g}^{-1}$ and $\sim 2000 \text{mAh cm}^{-3}$, respectively, which is much larger than the values of $\sim 372 \text{mAh g}^{-1}$ and $\sim 820 \text{mAh cm}^{-3}$ of graphite. In addition, $x$-graphdiyne would be advantageous over other 2D layers of sp-sp$^2$ hybrid carbon networks because the volume change is negligible through Li intercalation. These results suggest that multilayer $x$-graphdiyne has considerable potential as a promising high-capacity lithium ion battery anode material.

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