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First-Principles Study of Lithium and Sodium Atoms Intercalation in Fluorinated Graphite

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ABSTRACT The structure evolution of fluorinated graphite (CF*x*) upon the Li/Na intercalation has been studied by firstprinciples calculations. The Li/Na adsorption on single CF layer and intercalated into bulk CF have been calculated. The better cycling performance of Na intercalation into the CF cathode, comparing to that of Li intercalation, is attributed to the different strength and characteristics of the Li-F and Na-F interactions. The interactions between Li and F are stronger and more localized than those between Na and F. The strong and localized Coulomb attraction between Li and F atoms breaks the C—F bonds and pulls the F atoms away, and graphene sheets are formed upon Li intercalation.

KEYWORDS first-principles, Li/Na rechargeable batteries, fluorinated graphite.

1 Introduction

The use of fluorinated graphite (CF*x*) materials as the active cathode material in primary lithium batteries with the theoretical energy density reach to 2189 W h $kg⁻¹$ was first demonstrated by Watanabe et al. [1, 2]. A few years later, the Li/ CF_x batteries with energy density up to 560 W h $kg⁻¹$ were first commercialized by Matsushita Electric Industrial Co., Ltd. in Japan [3]. The high energy density is attributed to the large Gibbs free energy change of the Li intercalation reaction, which can be written as $(CF)_n + nLi \rightarrow (C)_n + nLiF$ [4]. In addition, the CF*x* has other advantages as electrode material such as high average operating voltage, long lifespan, stable operation, and high efficiency in primary lithium batteries [1, 2]. These advantages make it the superior cathode material in primary lithium batteries.

Recently, there are many attempts to extend the application of CF*x* in lithium ion batteries (LIBs). However, Li is easy to react with the fluorides and forms thermodynamically very stable LiF compound, as it is shown in the primary Li/ CF*x* batteries, and therefore the electrochemical reaction is

irreversible [5]. This makes it very difficult for the application of CF*x* as electrode material in LIBs, in which Li ions should be able to reversibly intercalate and extract [6]. Recently, Liu et al. reported that Na can be reversibly intercalated into and extracted from the CF*x* compound and the cycling of the Na-CF*x* batteries is good under room temperature [7]. The authors showed that the formation and decomposition of NaF are reversible in the electrochemical reaction, indicating that the structure of the CF_x is not destroyed or the structural change is reversible upon Na intercalation.

As Na and Li are belong to the same main group on the periodic table, it might be very interesting to know why Na can be reversibly intercalated/extracted into/from the CF*x*. What are the different characteristics between Li and Na that make the different performances upon Li and Na intercalation into the CF*x* compound? In the present work, we analyzed the structural evolution behavior of the CF*x* upon Na/Li intercalation by first-principles calculations.

2 Method and computational details

All calculations are performed using the Vienna Ab-initio Simulation Package (VASP) [8] within the projector augmentedwave (PAW) approach [9], with which the ion-electron interactions are described. The ground state of the electronic structure is described within the density functional theory (DFT) using the generalized gradient approximation (GGA) with PW91 exchange correlation functional [10]. We mention here that other functionals are also tested and no obvious difference is found in our calculations. The cutoff energy for expansion of the wave functions is 550 eV. The Monkhorst-Pack [11] scheme *k*-points sampling method is used for the integration in the first Brillouin zone and a 6 × 6 × 1 *k*-mesh is used in this study. The atomic positions are fully relaxed until the force on each atom is less than 0.02 eV \AA ⁻¹. We used a 3 \times 3 \times 1 CF supercell (including 36 C and 36 F atoms) to simulate the bulk phase of the chair configuration (the lowest en-

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ergy configuration) with AB stacking. The bulk phase of the CF with AA stacking was also tested, and the results showed that the total energy of the AA stacking is higher than that of the AB stacking, and therefore, all results presented in this work is based on the AB stacking model. The single CF layer model is obtained from the bulk model and the CF layers are separated by a 15 Å thick vacuum layer, in order to ensure that the interactions between layer and layer are negligible.

3 Results and discussion

3.1 Adsorption of Li/Na on single layer CF

In order to find out the interaction between the Li/Na and the CF layers, the single layer of CF with one Li/Na atom adsorption on it has been calculated. Different adsorption sites are tested and the stable adsorption site for single Li/Na atom is the hollow site. The optimized structures are shown in Figure 1. The distances from Li atom and Na atom to the nearest F atom (No. "1" in Figure 1, denoted as $F^{\text{\'et}}$) are 1.859 Å and 2.207 Å, respectively. As both the Li and Na ions are univalent cations, the Coulomb interactions between the Li/Na and F are mainly determined by the distance, and thus the Li-F interaction is stronger than the Na-F interaction in the CF layer.

Figure 1. The optimized structures with one Li/Na atom adsorption on single CF layer. The numbers from 1 to 6 denote different F atoms close to the adsorbate. (a) $C_{18}F_{18}$; (b) Li $C_{18}F_{18}$; (c) Na $C_{18}F_{18}$.

Then, we further analyzed the change of C—F bond lengths upon the adsorption of the Li/Na atom. The data are summarized in Table 1, from which we can see that the C—F bonds are elongated substantially upon Li/Na adsorption. As the $F^{\prime\prime}$ atom, as shown in Figure 1, is closest to the Li/Na atom, the corresponding C—F bond length becomes the longest due to the strongest Coulomb attraction. On the other hand, the C—F bond length in the case of Li adsorption (1.493 Å) is obviously longer than that in the case of Na adsorption (1.458 Å), indicating that the interaction between the Li and F"1" atoms is much stronger than that between the Na and $F^{\prime\prime\prime}$ atoms. The distance of Li-F and Na-F is one important reason for this, as discussed above, but it is not the only one. Our Bader charge [12] analysis also demonstrated that the Li is more ionized than Na upon adsorption on the CF layers. The Bader charges of Li and Na atoms adsorbed on the CF layer are 0.103*e* and 0.152*e*, respectively. That is to say, the charge transfer from Li to F is more than that from Na to F. Therefore, the stronger Li-F attraction is also contributed by the more charges they possess (Li is positively charged while F is negatively charged), comparing to that of the Na-F case.

Table 1. C—F bond lengths (unit: Å).

F site	$C_{18}F_{18}$	$LiG_{18}F_{18}$	$NaC_{18}F_{18}$	
n_1	1.383	1.493	1.458	
$^{\prime\prime}2^{\prime\prime}$	1.383	1.408	1.404	
"3"	1.383	1.405	1.410	
"4"	1.383	1.389	1.392	
"5"	1.383	1.400	1.404	
100''	1.383	1.394	1.395	

Very interestingly, the $C-F^{2^r}$ bond length is also slightly elongated upon the adsorption of Li and Na atoms, although the $F^{\prime 2}$ is located at the opposite side of the adsorbate. The reason is also related with the charge transfer from the Li/ Na atoms to the F atoms. In the CF layer, the electronic configuration of the C atom is in sp^2 hybridization, and the p_z electron is transferred to the F atom. Therefore, the planner hexatomic carbon ring is distorted and the buckling of the structure occurs. Upon Li/Na adsorption, charges transferred from Li/Na to F atom, and in turn the amount of the charges transferred from C to F becomes small, which also leads to a weakened C—F bonding. As a result, all of the C—F bond lengths become longer, as shown in Table 1. As it can be seen, the C—F bond lengths are sensitive to the distance between the adsorbate and the F atom. Furthermore, we can also see that this effect is more clearly around the Li atom, indicating that the charge distribution (F-Li interaction) is more localized for Li adsorption than that of Na adsorption.

In order to quantitatively describe the interactions between the adsorbate and F atoms, we calculated the binding energies defined as

$$
E_{\text{LiF}} = E(\text{Li}) + E(\text{F}) - E(\text{LiF})
$$

$$
E_{\text{LiC}_{18}\text{F}_{18}} = E(\text{Li}) + E(\text{C}_{18}\text{F}_{18}) - E(\text{LiC}_{18}\text{F}_{18})
$$

where the E_{Lif} and $E_{\text{Lic}_{18}F_{18}}$ are the binding energies of LiF molecule and Li on CF layers, respectively; the *E*(Li) is the total energy of LiF molecule in vacuum; $E(\text{LiC}_{18}F_{18})$ and $E(\text{C}_{18}F_{18})$ are the total energies of the single layer of CF with and without Li atom adsorption, respectively; *E*(Li) and *E*(F) are the total energies of the corresponding atom in vacuum. The definition for Na adsorption is the same as that of Li. The results are given in Table 2. As it can be seen, the binding energy of

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