


 Cite this: *RSC Adv.*, 2020, 10, 23573

Essential geometric and electronic properties in stage-*n* graphite alkali-metal-intercalation compounds

 Wei-Bang Li,^a Shih-Yang Lin,^b Ngoc Thanh Thuy Tran,^c Ming-Fa Lin ^{*ac} and Kuang-I Lin^d

The rich and unique properties of the stage-*n* graphite alkali-metal-intercalation compounds are fully investigated by first-principles calculations. According to the main features, the lithium and non-lithium (Na, K, Rb, Cs) systems are quite different from each other in stacking configurations, intercalant alkali-metal-atom concentrations, free conduction electron densities, atom-dominated and (carbon, alkali metal)-co-dominated energy bands, and interlayer charge density distributions. The close relations between the alkali-metal-doped metallic behaviors and the geometric symmetries are clarified through the interlayer atomic interactions. The stage-1 graphite alkali-metal-intercalation compounds possess the highest charge distribution for all stage-*n* types; moreover, those of the lithium systems are greater than those of the non-lithium systems. The lithium systems also have the largest blue shift of the Fermi level among all alkali metal systems.

Received 21st January 2020

Accepted 22nd April 2020

DOI: 10.1039/d0ra00639d

rsc.li/rsc-advances

Introduction

Graphene is a single layer with pure carbon honeycomb lattice.¹ Bulk graphite,² which could be regarded as a layered graphene system,^{3–6} has attracted a lot of theoretical and experimental research in basic science,^{4,6–8} engineering⁷ and applications.⁸ 3D graphite systems might present AA, AB, ABC⁹ and turbostratic stackings.¹⁰ All of them belong to semimetals under the interlayer hopping integrals of C-2p_z orbitals. In general, such condensed-matter systems become n- or p-type metals, depending on the kinds of intercalated atoms or molecules.¹¹ For example, the intercalation of alkali metal atoms¹² or FeCl₃ (ref. 13) into graphite creates many free conduction electrons and valence holes, as observed in a pure metal. Also, graphite could serve as the best anode material¹⁴ in commercial Li⁺-based batteries,¹⁵ mainly owing to the lowest cost, the most stable structure for intercalation, and the outstanding ion transport under charging and discharging processes. When the Li⁺ ions are released from the cathode, they will transport through the electrolyte, and then intercalate into the graphitic system. Most importantly, the flexible interlayer spacings between graphene layers are capable of providing sufficient

positions for the various intercalant concentrations.¹⁶ Any intermediate states and meta-stable configurations, which are created during the ion/atom intercalation, could survive through the very strong σ bondings in graphitic sheets. As a result, graphite is rather suitable for studying structural transformations in chemical reactions. For example, the close relations between graphene and intercalant layers in lattice symmetries are expected to present a dramatic transformation before and after the intercalation/de-intercalation processes.

Up to now, there have been a lot of theoretical and experimental studies on the fundamental properties in graphite-related systems. The former covers the phenomenological models and number simulation methods. For example, the tight-binding model, the random-phase approximation, and the Kubo formula are, respectively, utilized to investigate the electronic properties and magnetic quantization behaviors,¹⁷ Coulomb excitations¹⁸ and impurity screenings,¹⁹ and optical absorption spectra.²⁰ Furthermore, first-principles calculations²¹ are available for understanding the optimal stacking configurations, intercalant lattices, π , σ and intercalation-induced energy bands, free conduction electrons/valence holes, and density of states (DOS).

Previous studies of alkali-metal-intercalation graphite compounds have primarily focused on Li, Na and K.^{22–24} However, certain critical physical quantities and pictures are absent from the previous studies, such as the atom-dominated band structures, the spatial charge densities between intercalant and graphene layers, the interlayer orbital hybridization of intercalant and carbon atom, and the atom- and orbital-projected van Hove singularities.

^aDepartment of Physics, National Cheng Kung University, Tainan, Taiwan. E-mail: mflin@ncku.edu.tw

^bDepartment of Physics, National Chung Cheng University, Chiayi, Taiwan

^cHierarchical Green-Energy Materials (Hi-GEM) Research Center, National Cheng Kung University, Tainan, Taiwan

^dCenter for Micro/Nano Science and Technology, National Cheng Kung University, Tainan, Taiwan



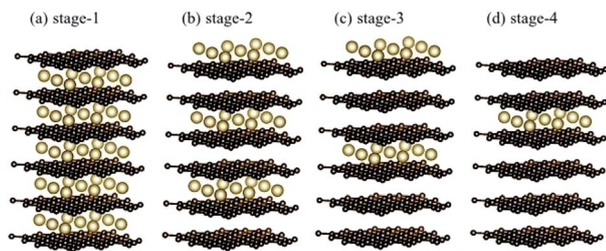


Fig. 1 The stage- n graphite alkali-metal-intercalation compounds: (a) $n = 1$, (b) $n = 2$, (c) $n = 3$, and (d) $n = 4$.

The main focuses of our work are the features of geometric and electronic properties in stage- n graphite alkali-metal-intercalation compounds (Li, Na, K, Rb, Cs) using the first-principles method.²¹ We aim to use first-principles calculation to understand the relation between the kinds of intercalants, concentrations and the essential properties, *e.g.*, lattice symmetries of intercalant layers, the dominances of valence and conduction states by carbon and/or alkali metal atoms, the blue shift of the Fermi level, the interlayer charge density variations after the alkali-metal intercalations, and the atom- and orbital-decomposed DOS.

Theoretical calculations

First-principles density functional theory (DFT)^{26–29} calculations are used to investigate the optimal geometric structures, charge distribution, and electronic, magnetic and optical characteristics of intercalation compounds. In this paper, we performed

the calculations with the Vienna *ab initio* simulation package (VASP), which evaluates an approximate solution within DFT by solving the Kohn–Sham equations. The Perdew–Burke–Ernzerhof formula used in VASP, which depends on the local electron density, is utilized to deal with many-particle Coulomb effects. As for the frequent electron-crystal scatterings, they are characterized by the projector-augmented wave pseudopotentials. The electron Bloch wave functions are solved from the linear superposition of plane waves, with a maximum kinetic energy of 500 eV. The current study on stage- n graphite compounds shows that the first Brillouin zone is sampled by $9 \times 9 \times 9$ and $100 \times 100 \times 100$ k -point meshes within the Monkhorst–Pack scheme, respectively, for the optimal geometry and band structure. Moreover, the convergence condition of the ground state energy is set to be $\sim 10^{-5}$ eV between two consecutive evaluation steps, where the maximum Hellmann–Feynman force for each ion is below 0.01 eV \AA^{-1} during the atom relaxations.²⁵

Results and discussion

(1) Unique stacking configurations and intercalant distributions

The normal stacking configurations, which are formed by layered graphene systems, include AAA, ABA, ABC and AAB. The interlayer spacings of graphitic layers are able to provide very suitable chemical environments for alkali metal intercalations and de-intercalations, *i.e.*, they are very useful during the charging and discharging processes. Chemical modifications hardly affect the planar honeycomb lattices and thus do not

Table 1 The stage-dependent ground state energies, the optimal geometric properties, the blue shifts of the Fermi levels, and the red shifts of the initial σ -electronic states for the stage- n graphite alkali-metal-intercalation compounds

	Stage-1, alkali-metal-doped graphite					Stage-2, alkali-metal-doped graphite				
	LiC ₆	NaC ₈	KC ₈	RbC ₈	CsC ₈	LiC ₁₂	NaC ₁₆	KC ₁₆	RbC ₁₆	CsC ₁₆
Interlayer distance with intercalation (Å)	3.815, 3.74, ³² 3.76 (ref. 30) (exp.)	4.601, 4.52 (ref. 32)	5.324, 5.40, ³² 5.35 (ref. 31) (exp.)	5.824, 5.65 (ref. 32)	6.032, 5.94 (ref. 32)					
Ground state energy (eV)	−57.29	−74.96	−75.13	−75.04	−75.12	−112.52	−148.78	−148.93	−148.84	−148.94
Blue shift of E_F in DOS	1.80	1.54	1.48	1.41	1.58	1.37	1.23	1.17	1.12	1.10
Blue shift of E_F in band structure (eV)	1.28–2.20	1.47–1.66	1.47–1.51	1.60–1.64	1.59–1.65	1.34–1.38	1.19–1.20	1.15–1.16	1.08–1.09	1.06–1.06
Red shifts of initial σ -electronic (eV)	1.25	1.24	1.14	1.13	1.0	1.22	0.99	0.93	0.86	0.82
	Stage-3, alkali-metal-doped graphite					Stage-4, alkali-metal-doped graphite				
	LiC ₁₈	NaC ₂₄	KC ₂₄	RbC ₂₄	CsC ₂₄	LiC ₂₄	NaC ₃₂	KC ₃₂	RbC ₃₂	CsC ₃₂
Interlayer distance with intercalation (Å)	3.864	4.878	5.360	6.056	6.058					
Ground state energy (eV)	−166.99	−222.64	−222.81	−222.69	−222.79	−223.15	−289.80	−296.55	−296.42	−296.48
Blue shift of E_F in DOS	1.07	1.01	0.98	0.94	0.90	0.89	0.83	0.80	0.77	0.78
Blue shift of E_F in band structure (eV)	1.22–1.23	1.02–1.03	0.57–0.58	0.84–0.84	0.76–0.77	0.31–0.33	0.63–0.64	0.63–0.63	0.51–0.52	0.41–0.41
Red shifts of initial σ -electronic (eV)	1.21	1.02	0.96	0.66	0.60	1.23	0.90	0.88	0.70	0.58



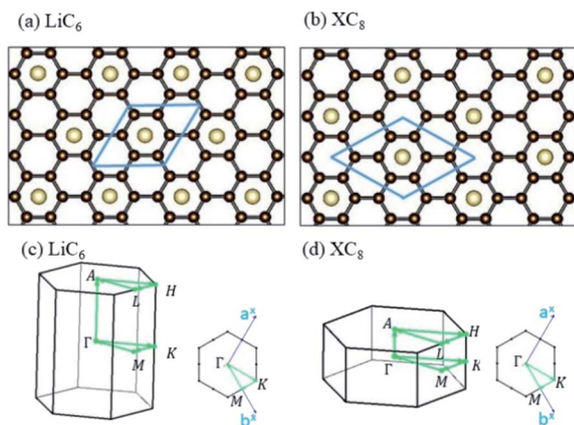


Fig. 2 The planar structures of (a) LiC_6 and (b) XC_8 [$X = \text{Na, K, Rb}$ and Cs]. The first Brillouin zones of (c) LiC_6 and (d) XC_8 [$X = \text{Na, K, Rb}$ and Cs].

change the orthogonal feature of π and σ bondings. Very interesting, the optimal geometric properties are rather sensitive to the (x, y) -plane distribution and concentration of alkali metal atoms, *e.g.*, the interlayer distances of neighboring planes and an enlarged primitive unit cell. According to the highest concentration, the graphite alkali-metal-intercalation materials could be classified into two categories: lithium and non-lithium ones. There exist LiC_{6n} or AC_{8n} for the stage- n materials [Fig. 1(a)–(d)], as examined from experimental measurements and the theoretical predictions. The specific distance between carbon and intercalant layer is shortest/longest for Li/Cs [Table 1], only directly reflecting the effective atomic radius. Although all the alkali metal atoms present hollow-site optimal positions, their distribution symmetries are dominated by Li atoms or non-Li ones. The layered LiC_{6n} and AC_{8n} systems, respectively, possess a three- and four-times enhancement in the unit cell [Fig. 2(a) and (b)]; therefore, their reduced first Brillouin zones exhibit different high-symmetry points [Fig. 3(a)]. This is expected to have a strong effect on the initial π -electronic states of the Dirac-cone structure. More chemical bonds, which arise from the alkali metal and carbon atoms, are produced in graphite intercalation compounds compared with pristine systems. Such bondings need to be taken into account for the diverse essential properties. In addition to the intercalant distribution, the stacking configuration of two neighboring graphitic sheets also affects the ground state energy. The current study clearly shows that only the stage-1 LiC_6 presents AA stacking, and the other compounds exhibit AB stacking. That is to say, all the graphite alkali-metal-intercalation materials exhibit the AB-stacked configurations except for the former system.

(2) Metallic and semi-metallic behaviors

Without alkali metal atom intercalations, pristine graphite exhibits unique semi-metallic behavior through weak, but significant van der Waals interactions. A semiconducting zero-gap graphene is dramatically changed into a semi-metallic

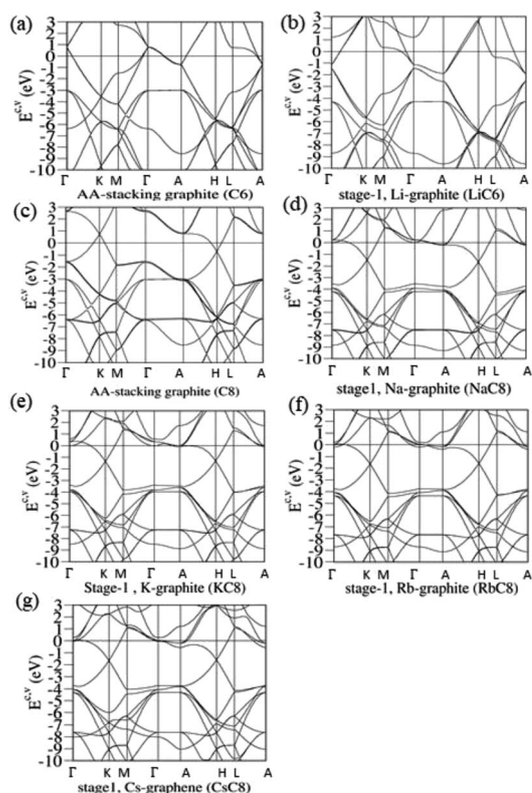


Fig. 3 The rich band structures of the pristine graphite and stage-1 graphite alkali-metal-intercalation compounds: (a) AA, (b) LiC_6 , (c) AB stackings without intercalations, (d) NaC_8 , (e) KC_8 , (f) RbC_8 and (g) CsC_8 .

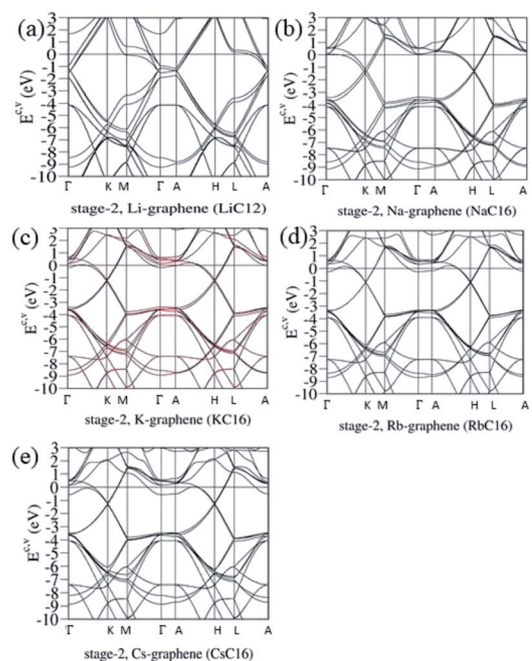


Fig. 4 The unusual electronic energy spectra of the AB-stacked stage-2 graphite alkali-metal-intercalation compounds: (a) LiC_{12} , (b) NaC_{16} , (c) KC_{16} , (d) RbC_{16} , and (e) CsC_{16} .



system for any few-layer stackings, or infinite-layer graphites. For example, there exist the unusual overlaps of valence and conduction bands along the KH path [Fig. 3(c); or the Γ K path under the reduced Brillouin zone in Fig. 3(b)] in AA-, AB- and ABC-stacked graphites through single-orbital interlayer atomic interactions. According to the VASP, the simple hexagonal/rhombohedral graphite [Fig. 3(b)], with the highest/lowest stacking symmetry, presents the largest/lowest free carrier density of conduction electrons and valence holes. In general, the π valence bands, which are purely related to C-2p_z, show the initial states at the stable K valley [Fig. 3(c)], the saddle-point structure near the M point [$E^v \sim -3$ eV], and its termination in the Γ valley [the whole π -band width more than 7 eV].

The band structures of lithium- and non-lithium graphite intercalation compounds sharply contrast with each other, such as those of stage-1 LiC₆ and XC₈ compounds in Fig. 3(d) and (e)–(h), respectively. For the former, the Fermi level is transferred from the middle of Dirac-cone structures into the conduction ones near the Γ valley after the intercalation of lithium atoms. That is to say, E_F presents a blue shift. Apparently, the free carriers are due to the outmost 2s orbitals of lithium atoms. The occupied and unoccupied states are highly asymmetric to each other about the Fermi level. The initial π/π^* valence/conduction states are mainly determined by the stacking configurations in the chemical intercalations [the distribution symmetry of Li-intercalants in Fig. 2(a)], or the corresponding relation between the original and reduced first Brillouin zone. There exists an observable energy spacing of $E^c \sim 0.31$ – 0.63 eV along the Γ –A path in the modified valence and conduction Dirac-cone structures. The creation of discontinuous states in

honeycomb lattices might arise from the different ionization energies of Li-2s and C-2p_z orbitals [the distinct on-site Coulomb potential energies]. The whole π valence-band width of LiC₆, being created by the C-2p_z orbitals, could be identified to about 7.32 eV/7.40 eV from the electronic energy spectrum along the Γ –M–K– Γ /A–H–L–A or Γ –K–M– Γ /A–L–H–A paths. This result clearly illustrates the well-behaved π bondings in graphite alkali-metal-intercalation compounds. Other σ bondings behave so, as indicated from the initial states at the Γ point of $E^v \sim -4.20$ eV. Their orthogonal relation remains the same after chemical modification, being very useful in establishing the tight-binding model for graphite alkali-metal-intercalation compounds.

The main features of the band structure, as clearly indicated in Fig. 3(e)–(h), are dramatically changed for the other alkali-metal-atom intercalations. Both stage-1 AC₈ and LiC₆ have the totally different distribution configurations [Fig. 2(a) and (b)], and so do the reduced first Brillouin zones [Fig. 3(a)]. For the former, the Dirac-cone structures of the π and π^* bands are initiated from the stable K/K' valleys, but not the Γ ones. The energy spacing of separated Dirac points is small or almost vanishing; furthermore, the Fermi level is situated above the conduction point about ~ 1.35 – 1.50 eV. In addition to E_F , whether or not the second conduction energy subband is partially occupied depends on the kinds of alkali metal atoms, such as the alkali-metal-induced free electrons in the first and second subbands for KC₈/RbC₈/CsC₈ [Fig. 3(f)/(g)/(h)]. The low-energy bands belong to the single states, without the split double degeneracy [Fig. 3(d) for LiC₆]. The whole π -band energy spectra are identified from the K– Γ –K–M– Γ and H–A–H–L–A

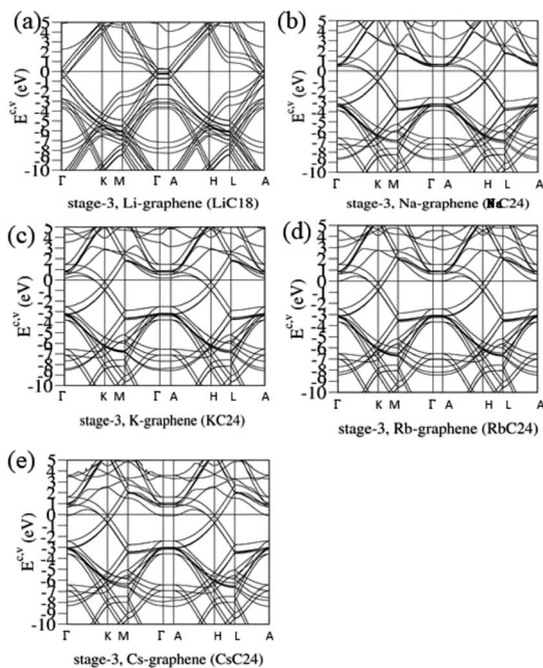


Fig. 5 The rich band structures for the AB-stacked stage-3 graphite alkali-metal-intercalation compounds: (a) LiC₁₈, (b) NaC₂₄, (c) KC₂₄, (d) RbC₂₄, and (e) CsC₂₄.

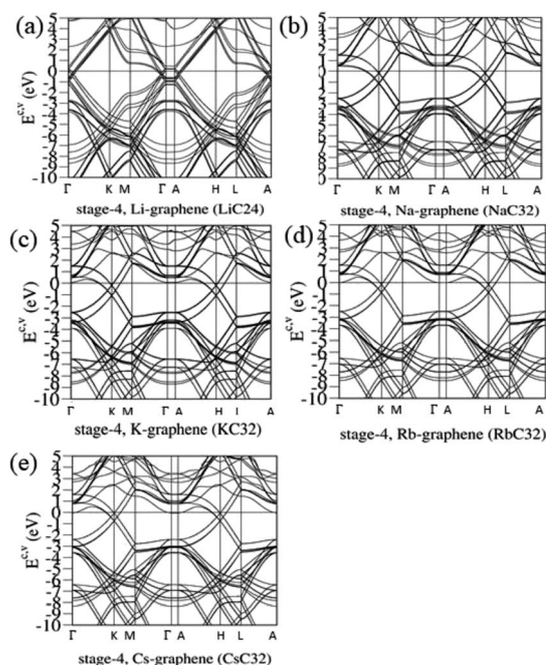


Fig. 6 The rich band structures for the AB-stacked stage-4 graphite alkali-metal-intercalation compounds: (a) LiC₂₄, (b) NaC₃₂, (c) KC₃₂, (d) RbC₃₂, and (e) CsC₃₂.



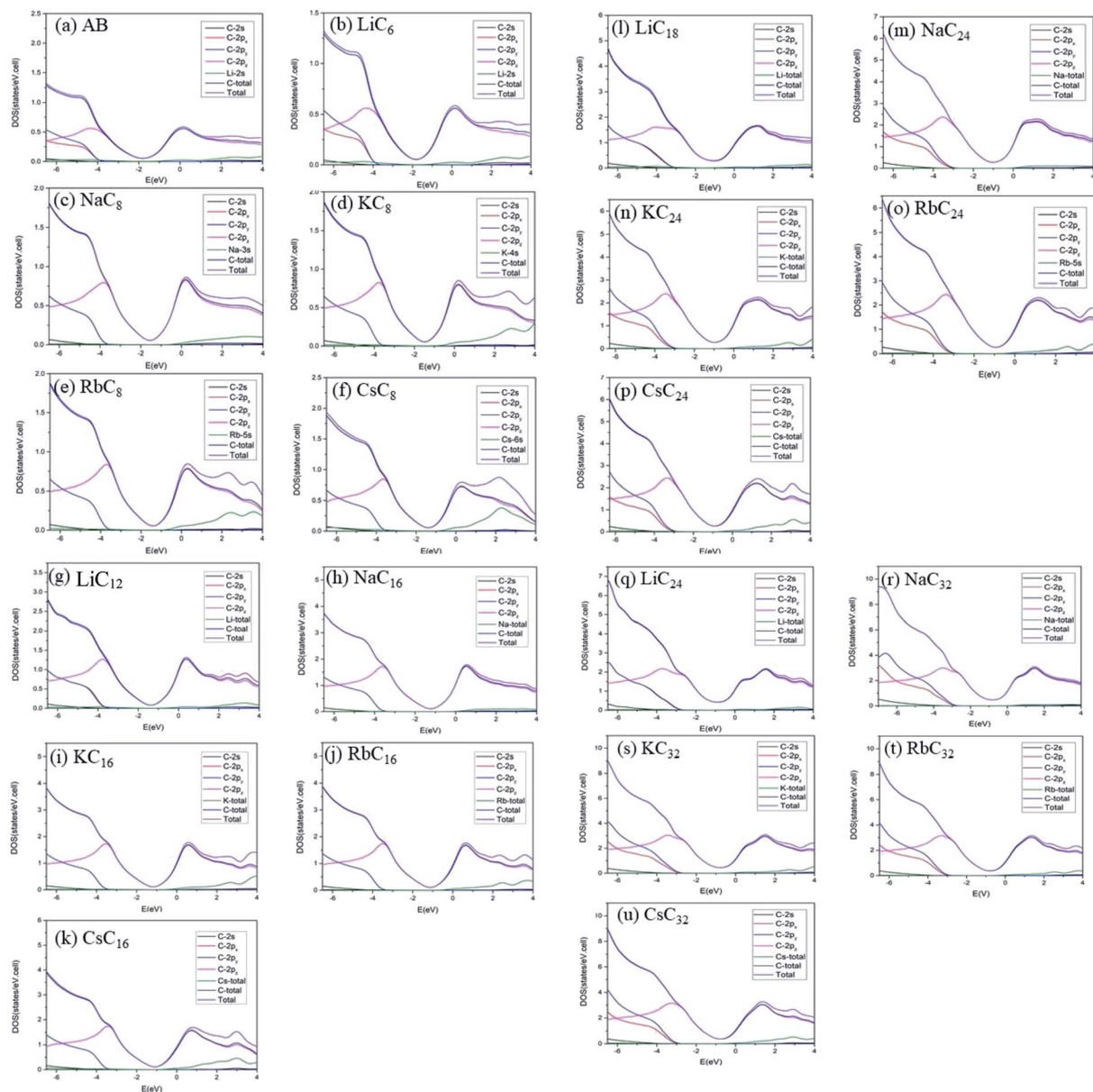


Fig. 7 The atom- and orbital-decomposed density of states for (a) AA, (b) LiC_6 , (c) NaC_8 , (d) KC_8 , (e) RbC_8 , (f) CsC_8 , (g) LiC_{12} , (h) NaC_{16} , (i) KC_{16} , (j) RbC_{16} , and (k) CsC_{16} , (l) LiC_{18} , (m) NaC_{24} , (n) KC_{24} , (o) RbC_{24} , (p) CsC_{24} , (q) LiC_{24} , (r) NaC_{32} , (s) KC_{32} , (t) RbC_{32} , (u) CsC_{32} . The black, red, light blue, pink, green, deep blue, and purple curves represent, respectively, C-2s, C-2p_x, C-2p_y, C-2p_z, alkali metal, carbon, and compound.

paths, leading to a width of $\sim 7.31\text{--}7.41$ eV. Very interesting, one pair of π -valence subbands come to exist near $E^v \sim -4.0$ eV, being accompanied by the initial pair of σ valence bands. This further illustrates the zone-folding effects on band structure and the well separation of π and σ bondings.

Electronic energy spectra strongly depend on the n stage of graphite intercalation compounds, as clearly illustrated in Fig. 4(a)–(e). Stage-2 LiC_{12} and AC_{16} have different first Brillouin zones on the (k_x, k_y) plane, and their k_z -ranges are associated with the distances between two intercalant planes [Fig. 1]. Compared with stage-1 band structures [Fig. 3(d)–(g)], the number of Dirac cones becomes double in the stage-2 cases, in which the further modifications include the reduced blue shift

of the Fermi level, the enhanced anisotropy, the induced energy spacings between valence and conduction Dirac points, and the diverse energy relations near the band-edge states. For example, four/two valence and conduction pairs arise from the Γ/K valley [or the A/H valley] for $\text{LiC}_{12}/\text{AC}_{16}$ [Fig. 4(a) and (b)–(e)]. Furthermore, the whole π -band widths could be roughly estimated from the $\Gamma\text{KM}\Gamma/\text{KMK}\Gamma$ path [or the ALHA/HLHA path]. The low-energy essential properties are dominated by the π bondings of C-2p_z orbitals. The above-mentioned obvious changes directly reflect the great enhancement of the interlayer atomic interactions due to the C–C bonds in two neighboring graphitic sheets/graphene-intercalant layers. Very interesting, all the stage-2 compounds exhibit a pair of σ bands at the Γ and



A valleys, and the energy dispersions along ΓA are negligible. These results indicate the mutual orthogonality of the planar σ and perpendicular π bondings. Such a phenomenon is expected to survive in any stage- n graphite alkali-metal-intercalation compounds.

The blue shifts about the Fermi level and the free conduction electrons decline quickly with an increase of the n number. The concentration of alkali metal atoms is greatly reduced, and so is the charge transfer from their outmost s -orbitals to carbon $2p_z$ -orbitals. Such results are clearly revealed in the stage-3 systems. Fig. 5 shows the slight modifications of the Dirac-cone structures and the intersecting of the Fermi level with most of the conduction bands. LiC_{18} [Fig. 5(a)] and AC_{24} [Fig. 5(b)–(e)], respectively, possess six and three pairs of linear valence and conduction bands near the Γ/A and K/H valleys. Apparently, there are significant changes in the observable energy spacing between valence and conduction Dirac points, the anisotropic Fermi velocities, and the distinct Fermi momenta slopes. The blue shifts of E_F are estimated to be 0.45, 0.58, 0.55, 0.53, and 0.5 eVs. Very interesting, the electronic energy spectra become more complicated under the stronger zone-folding effects. The main features of electronic properties are closely related to the weak, but important Li–C bonds, *e.g.*, the minor contributions of alkali metal atoms on each electronic state. That is, the diversified n -type dopings are created by the critical Li–C and C–C bondings.

According to the spatial charge distributions and their variations after chemical modifications [Fig. 8–12], the chemical bondings hardly depend on the distributions and concentrations of alkali metal adatoms. After the alkali-metal-atom intercalations, we can observe that the changes depend on the

concentrations and the kinds of alkali metal atoms. Comparing the charge distributions ρ with the charge variation $\Delta\rho$, we can find the orbital hybridization in A–C bonds ($A = \text{Li}, \text{Na}, \text{K}, \text{Rb}, \text{Cs}$), and the charge density almost remains the same in the graphitic layers without direct intercalation with the alkali metal atoms, *e.g.*, the middle one layer/two layers in stage-3/stage-4 compounds. Moreover, with increasing n number, $\Delta\rho$ becomes small [$\Delta\rho$: stage-1 > stage-2 > stage-3 > stage-4]. We also see the evidence of $2s$ – $2p_z$ hybridization in A–C bonds in the charge variation. Obviously, the stage-1 compounds exhibit the strongest electron transfer from alkali metal atoms to carbons, as shown by the red region in $\Delta\rho$ [Fig. 8–12]. The variation reduces when the radius of alkali metal atoms gets larger, that is to say the electron transfer from A to carbon atoms is strongest in Li but weakest in Cs [radius: $\text{Li} < \text{Na} < \text{K} < \text{Rb} < \text{Cs}$; variation: $\text{Li} > \text{Na} > \text{K} > \text{Rb} > \text{Cs}$].

The atom- and orbital-dependent DOSs [Fig. 6] allow a full understanding of the metallic behavior and the close relations among the different chemical bondings. Although band structures become very complicated under the alkali-metal-atom intercalations, the main features of van Hove singularities are sufficiently clear for the identifications of diverse phenomena through a suitable broadening factor. The low-energy DOSs in stage-1 and stage-2 graphite alkali-metal-intercalation compounds present a prominent peak just at the Fermi level [$E_F = 0$], regardless of the kind of alkali metal atoms. Furthermore, there exists a valley structure, with a minimum value, at its left-hand neighbor. When such a characteristic is combined with the similar ones at E_F , the Fermi level is deduced to exhibit a blue shift. That is to say, E_F is situated at the conduction

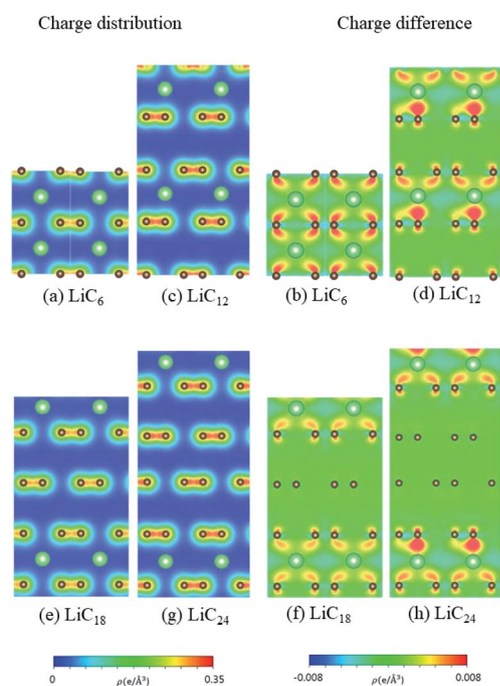


Fig. 8 The spatial charge distribution before/after Li intercalation: (a and b) LiC_6 , (c and d) LiC_{12} , (e and f) LiC_{18} , (g and h) LiC_{24} .

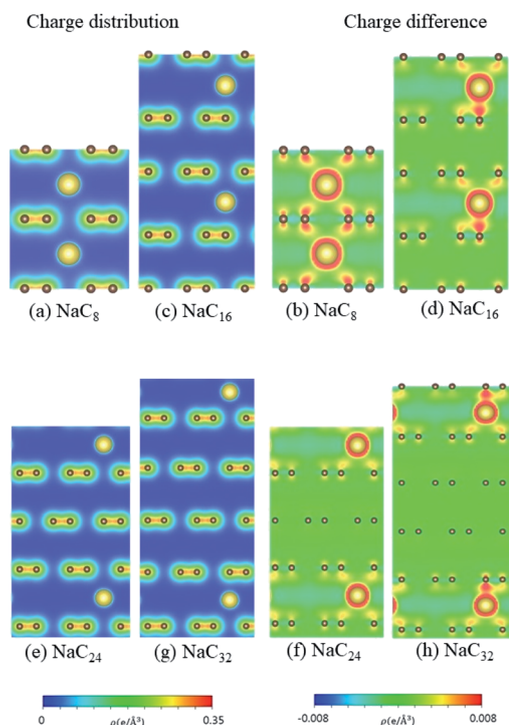


Fig. 9 The spatial charge distribution before/after Na intercalation: (a and b) NaC_8 , (c and d) NaC_{16} , (e and f) NaC_{24} , (g and h) NaC_{32} .



energy subbands [E_F roughly lies in the center of valence and conduction bands] roughly after [before] the alkali-metal-atom intercalations. It should be noticed that the contributions due to the alkali metal atoms are weak, but rather important. The blue shifts of stage-1 LiC_6 , NaC_8 , KC_8 , RbC_8 and CsC_8 , and stage-2 LiC_{12} , NaC_{16} , KC_{16} , RbC_{16} and CsC_{16} are, respectively, estimated to be 1.80, 1.54, 1.48, 1.41, 1.58, 1.37, 1.23, 1.17, 1.12, 1.10 eV.

The van Hove singularities in graphite intercalation compounds [Fig. 7(b)–(u)], which survive in the specific energy ranges, are mainly determined by the carbon or alkali metal atoms and their orbitals. Most important, DOSs in the critical energy range of $-5.0 \text{ eV} < E < 3.0 \text{ eV}$, being relatively easily examined from experimental STS measurements, are dominated by the C- $2p_z$ orbitals [the pink curves]. Furthermore, the minor contributions related to the outmost s-orbital of alkali metal atoms [the green curves], especially in the conduction energy spectrum, play a critical role in determining the blue shift of the Fermi level. As for the C- $[2p_x, 2p_y]$ orbitals [the red and light blue curves], their contributions are initiated from $\sim E < -4.0 \text{ eV}$, while they are absent in the opposite energy range. The red shift of the initial σ valence bands is about 1.0 eV, compared with those of the pristine simple hexagonal and Bernal graphites [Fig. 7(a)]. Only the LiC_6 case [Fig. 7(b)] exhibits the split contributions of $2p_x$ and $2p_y$ orbitals. This result directly reflects the anisotropic distribution configuration. Also, the C- $2s$ orbitals come to exist at the deeper energies of $\sim E < -5.0 \text{ eV}$. Apparently, the above-mentioned features indicate the

good separation of C- $2p_z$ and C- $[2s, 2p_x, 2p_y]$ orbital contributions, and thus the normal perpendicular orbital hybridizations of π and σ chemical bonds. That such bonding behavior is strongly linked with the significant interlayer $2p_z$ -orbitals due to the carbon-alkali metal bonds can account for the featured electronic properties, *e.g.*, the main features of band structures and DOSs in Fig. 3–7. The clear identifications of stage-1 and stage-2 of graphite alkali-metal-intercalation compounds could be achieved from the low-energy features of van Hove singularities, as indicated in Fig. 7(b)–(f) and (g)–(k), respectively. Compared with those of the former, the blue shifts of the Fermi levels are relatively small, in which they are, respectively, $\sim 1.20 \text{ eV}$, 1.15 eV , 1.10 eV , 1.06 eV and 1.03 eV for LiC_{12} , NaC_{16} , KC_{16} , RbC_{16} , and CsC_{16} . Most important, their DOSs at E_F do not belong to the local maxima. Such a result directly reflects whether or not the band-edge states of conduction bands are somewhat away from the Fermi level [Fig. 4(b)–(e)]. This significant difference between stage-2 and stage-1 systems further illustrates the intersecting of E_F and conduction bands, thus leading to the n-type doping cases after the alkali-metal-atom intercalations. Previous researches mainly show the structures, total energy, enthalpy of formations and the DOS.^{22–24,31–33} The geometric structures, especially the interlayer distances, are slightly different between these papers and experiment; for instance the LiC_6 distance is $3.73/3.76 \text{ \AA}$ in ref. 24/ref. 23, 3.81 \AA in this paper and 3.76 \AA in experiment³⁰ [other previously reported details in Table 1]. The results of our theoretical calculations are close to those of previous studies; furthermore, we give the details of geometric structures, ground

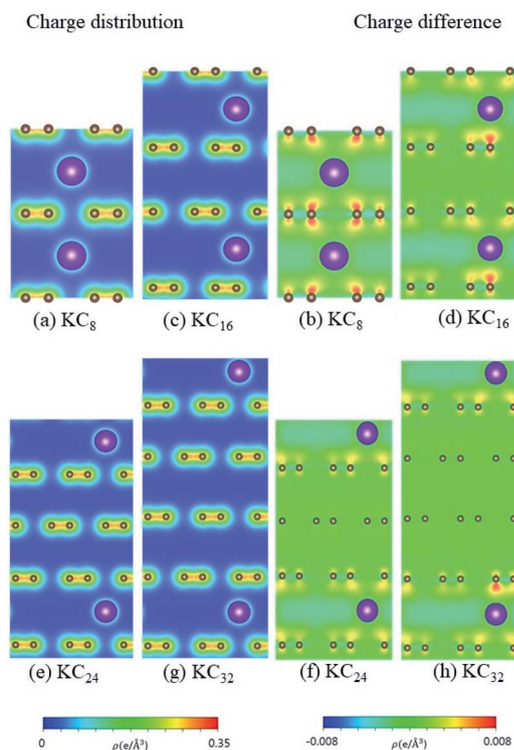


Fig. 10 The spatial charge distribution before/after K intercalation: (a and b) KC_8 , (c and d) KC_{16} , (e and f) KC_{24} , (g and h) KC_{32} .

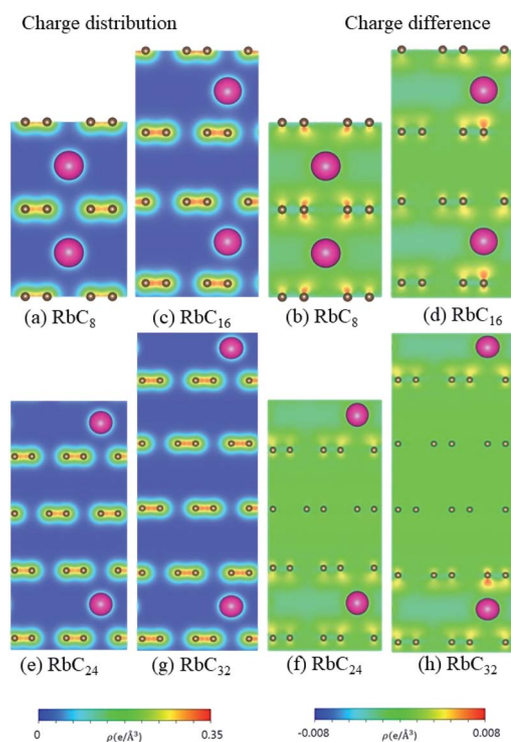


Fig. 11 The spatial charge distribution before/after Rb intercalation: (a and b) RbC_8 , (c and d) RbC_{16} , (e and f) RbC_{24} , (g and h) RbC_{32} .



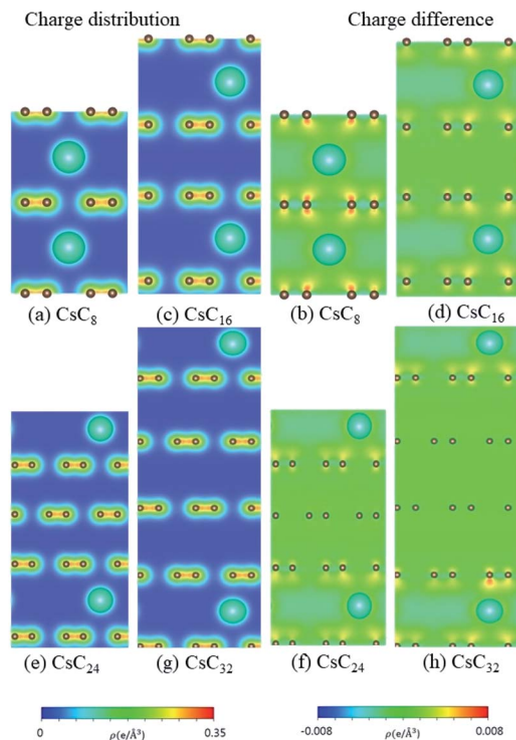


Fig. 12 The spatial charge distribution before/after Cs intercalation: (a and b) CsC_8 , (c and d) CsC_{16} , (e and f) CsC_{24} , (g and h) CsC_{32} .

state energies, blue shift (both in band structures and DOS) and spatial charge distributions.

Conclusion

The fundamental properties of (Li, Na, K, Rb, Cs)-intercalated graphite compounds under the distinct stage configurations have been investigated by means of first-principles calculations. Due to the band structures, DOS and spatial charge distributions [Fig. 3–12], the weak but significant van der Waals interactions, which arise from the interlayer $2p_z$ - $2p_z$ and $2p_z$ - s orbital hybridizations in C–C and C–A bonds, respectively, make the most important contribution to the low-lying π -electronic structure and thus dominate the essential physical properties. The dramatic changes include the blue shift of the Fermi level/the red shift of the σ bands [the n-type doping behaviors], the greatly enhanced asymmetric electron and hole energy spectra, and the obviously reduced conduction electron density for the dilute intercalant cases. These calculated results suggest that the essential electronic properties are relevant to the concentrations of intercalant alkali metal atoms. The radii of the distinct alkali metal atoms cause the very different interlayer distances, e.g., 3.8/4.3/5.3/5.8/6.0 Å for stage-1 Li/Na/K/Rb/Cs, respectively [Table 1]. Therefore, the smallest alkali metal atom, Li, must have the strongest orbital hybridization between the alkali metal atom and carbons. The charge transfers from the alkali metal to carbons decrease quickly with the increase of the n number, and the lithium systems possess the highest ratio

of charge transfer. The lithium systems also have the largest blue shift of the Fermi level of ~ 1.8 eV.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work is supported by the Hi-GEM Research Center and the Taiwan Ministry of Science and Technology under grant numbers MOST 108-2212-M-006-022-MY3 and MOST 108-3017-F-006-003.

References

- 1 M. Matthew and B. Regan, *Phys. Rev. Lett.*, 2011, **106**, 116803; M. Rafique, N. H. Mirjat, A. M. Soomro, S. Khokhar and Y. Shuai, *Phys. Lett. A*, 2018, **16**, 1108–1119.
- 2 S.-M. Lee, D.-S. Kang and J.-S. Roh, *Carbon Lett.*, 2015, **16**, 135–146.
- 3 N. T. T. Tran, S. Y. Lin, C. Y. Lin and M. F. Lin, *Geometric and Electronics Properties of Graphene Related Systems*, CRC Press, 2017, ISBN 9781351368483.
- 4 G. Gómez-Santos, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2009, **80**, 245424.
- 5 D. D. L. Chung, *J. Mater. Sci.*, 2002, **37**, 1475–1489.
- 6 Y. Wang, J. E. Panzik, B. Kiefer and K. K. Lee, *Sci. Rep.*, 2012, **2**, 520; J. H. Ho, C. P. Chang and M. F. Lin, *Phys. Lett. A*, 2006, **352**, 446–450; C. Y. Lin, R. B. Chen, Y. H. Ho and M. F. Lin, *Electronic and Optical Properties of Graphite-Related Systems*, CRC Press, 2017, ISBN 9781138571068.
- 7 Z. Zhang, H. Huang, X. Yang and L. Zang, *J. Phys. Chem. Lett.*, 2011, **2**, 2897–2905.
- 8 Q. Cheng, *et al.*, *Sci. Rep.*, 2017, **7**, 14782.
- 9 C.-Y. Lin, J.-Y. Wu, C.-W. Chiu and M.-F. Lin, *Work function of alkali-atom adsorbed graphite*, CRC Press, 2019, ISBN: 9780429277368.
- 10 M. Aoki and H. Amawashi, *Solid State Commun.*, 2007, **142**, 123–127.
- 11 R. Matsumoto, Y. Okabe and N. Akuzawa, *J. Electron. Mater.*, 2015, **44**, 399–406; X. Meng, S. Tongay, J. Kang, Z. Chen, F. Wu, S.-S. Li, J.-B. Xia, J. Li and J. Wu, *Carbon*, 2013, **53**, 507–514.
- 12 A. Natori, T. Ohno and A. Oshiyama, *J. Phys. Soc. Jpn.*, 1985, **54**, 3042–3050.
- 13 Y. Li and Y. Qu, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2013, **425**, 72–77.
- 14 C. Mao, *et al.*, Selecting the Best Graphite for Long-Life, High-Energy Li-Ion Batteries, *J. Electrochem. Soc.*, 2018, **165**, A1837–A1845.
- 15 Q. Cheng, *Sci. Rep.*, 2017, **7**, 14782; S. Byun, J. Park, W. A. Appiah, M.-H. Ryou and Y. M. Lee, *RSC Adv.*, 2017, **7**, 10915; M. Steinhauer, S. Risse, N. Wagner and K. Andreas Friedrich, *Electrochim. Acta*, 2017, **228**, 652–658; F. M. Kindermann, P. J. Osswald, S. Klink, G. Ehlert, J. Schuster, A. Noel, S. V. Erhard, W. Schuhmann and



- A. Jossen, *J. Power Sources*, 2017, **342**, 638–642; M. Bauer, B. Rieger, S. Schindler, P. Keil, M. Wachtler, M. A. Danzer and A. Jossen, *J. Energy Storage*, 2017, **10**, 1–10; H. P. T. Sasanka Hewathilake, N. Karunarathne, A. Wijayasinghe and N. W. B. Balasooriya, *Ionics*, 2017, **23**, 1417–1422.
- 16 S.-Y. Lin, N. T. T. Tran, S.-L. Chang, W.-P. Su and M. F. Lin, *Structure- and Adatom-Enriched Essential Properties of Graphene Nanoribbons*, CRC Press, 2018, ISBN: 9780367002299.
- 17 J. H. Ho, Y. H. Lai, S. J. Tsai, J. Hwang, C. Chang and M. F. Lin, *J. Phys. Soc. Jpn.*, 2006, **75**, 72–77; Y. K. Huang, S. C. Chen, Y. H. Ho, C. Y. Lin and M. F. Lin, *Sci. Rep.*, 2014, **4**, 7509; M. Koshino and E. McCann, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2011, **83**, 165443; Z. F. Wang, F. Liu and M. Y. Chou, *Nano Lett.*, 2012, **12**, 3833–3838.
- 18 J. H. Ho, C. P. Chang and M. F. Lin, *Phys. Lett. A*, 2006, **352**, 446–450; M. F. Lin, Y. C. Chuang and J. Y. Wu, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2012, **86**, 125434; J. Y. Wu, S. C. Chen, O. Roslyak, G. Gumbs and M. F. Lin, *ACS Nano*, 2011, **5**, 1026–1032; Y. E. Lozovik and A. A. Sokolik, *Nanoscale Res. Lett.*, 2012, **7**, 134; T. Ohta, *et al.*, *Phys. Rev. Lett.*, 2007, **98**, 206802.
- 19 L. M. Roth and G. W. Pratt Jr, *J. Phys. Chem. Solids*, 1959, **8**, 47–49; K. Schrüfer, C. Metzner, M. C. Hofmann and G. H. Döhler, *Superlattices Microstruct.*, 1997, **21**, 223–230; Z. L. Mišković, P. Sharma and F. O. Goodman, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2012, **86**, 115437.
- 20 Q. T. Ain, A. Al-Modlej, A. Alshammari and M. N. Anjum, *Mater. Res. Express*, 2018, **5**, 035017.
- 21 O. Leenaerts, B. Partoens and F. M. Peeters, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2008, **77**, 125416.
- 22 E. Ziambaras, J. Kleis, E. Schröder and P. Hyldgaard, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2007, **76**, 155425.
- 23 K. Persson, Y. Hinuma, Y. S. Meng, A. Van der Ven and G. Ceder, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2010, **82**, 125416.
- 24 Z. Wang, M. S. Sverre and T. Grande, *RSC Adv.*, 2014, **4**, 4069.
- 25 S.-Y. Lin, *et al.*, *Phys. Chem. Chem. Phys.*, 2015, **39**, 26443–26450.
- 26 M. Rafique, M. A. Unar, I. Ahmed, A. R. Chachar and Y. Shuai, *J. Phys. Chem. Solids*, 2018, **118**, 114–125.
- 27 M. Rafique, Y. Shuaia, H.-P. Tan and M. Hassan, *Appl. Surf. Sci.*, 2017, **408**, 21–33.
- 28 G. Kresse and J. Hafner, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1993, **47**, 558–561.
- 29 G. Kresse and J. Hafner, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1994, **49**, 14251–14269.
- 30 R. Juza and V. Wehle, *Naturwissenschaften*, 1965, **52**, 560.
- 31 E. Ziambaras, J. Kleis, E. Schröder and P. Hyldgaard, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2007, **76**, 155425.
- 32 G. E. Pascal and F. Mireille, *Solid State Ionics*, 1988, **28–30**, 1172–1175.
- 33 G. Yoon, H. Kim, I. Park and K. Kang, *Adv. Energy Mater.*, 2017, **7**, 1601519.

