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## 1. Introduction

Searching for the maximum coordination number in planar species has fascinated chemists for many years and continuously pushed the boundary of our understanding of chemical structures and bonding.<sup>1,2</sup> The central atom and periphery atoms around it in the ligand in stable planar hypercoordinated structures must match both geometrically and electronically, *i.e.*, they must have the right atomic sizes and electronic configurations. The experimentally observed boron-centered monocyclic boron wheel clusters  $D_{7h}$  B ${}^{\odot}B_{7}^{2-}$  and  $D_{8h}$  B ${}^{\odot}B_{8}^{--}$ are good examples in which the central B atoms have the coordination numbers of CN = 7 and 8, respectively.<sup>3</sup> These boron cluster monoanions prove to be  $\sigma + \pi$  dually aromatic in nature with six delocalized  $\sigma$  and six delocalized  $\pi$  electrons (6  $\sigma$ + 6  $\pi$ ) each conforming to the 4N + 2 aromatic rule ( $N_{\sigma} = N_{\pi} =$ 1). Based on the double aromaticity requirement in bare boron wheel clusters, a general electronic design principle x + n + k =12 or 16 was developed for metal-centered monocyclic boron wheel clusters  $M \otimes B_n^{k-}$  by the groups of Wang and Boldyrev,<sup>1,4-6</sup> where *x* stands for the formal valence of the metal center M. This design principle has been successfully applied to the

# Metal-centered monocyclic carbon wheel clusters with record coordination numbers in planar species†

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The highest coordination number identified to date in planar species is CN = 10 in metal-centered monocyclic boron wheel clusters  $D_{10h} \ M \otimes B_{10}^{-}$  (M = Ta and Nb) (Galeev *et. al., Angew. Chem. Int. Ed.,* 2012, **51**, 2101). Extensive global minimum searches and first-principles theory calculations performed herein indicate that the experimentally observed  $LaC_{13}^{+}$  and  $LaC_{14}^{+}$  possess the well-defined global minima of perfect metal-centered monocyclic carbon wheel  $D_{13h} \ La \otimes C_{13}^{+}$  (1)  $(1A'_1)$  and slightly off-centered  $C_{2v} \ La \otimes C_{14}^{+}$  (4) (<sup>1</sup>A<sub>1</sub>) with record coordination numbers of CN = 13 and 11 in planar structures, respectively, further pushing the boundary of our understanding of chemical structures and bonding. Detailed molecular orbital, nucleus-independent chemical shift, and ring current analyses indicate that  $D_{13h} \ La \otimes C_{13}^{+}$  (1) is  $\sigma + \pi$  dually aromatic in nature, with 14 totally delocalized in-plane  $\sigma$  electrons and 14 totally delocalized out-of-plane  $\pi$  electrons each matching the 4N + 2 aromatic rule ( $N_{\sigma} = N_{\pi} = 3$ ). Similar  $\sigma + \pi$  dually aromatic metal-centered monocyclic wheel clusters  $D_{13h} \ Ca \otimes C_{13}$  (2),  $C_{13v} \ Ac \otimes C_{13}^{+}$  (3),  $C_{2v} \ Y \otimes B_6 C_6^{+}$  (5), and  $C_{2v} \ S C \otimes B_5 C_6$  (6) have also been obtained with CN = 13, 13, 12, and 11, respectively. The results obtained in this work effectively enrich the chemical structures and bonding patterns of planar hypercoordinated complexes.

experimentally characterized octacoordinated  $D_{8h}$  Co $@B_8^-$  with CN = 8,<sup>5</sup> nonacoordianted  $D_{9h}$  Ru $@B_9^-$ ,  $D_{9h}$  Rh $@B_9^-$ , and  $D_{9h}$  Ir $@B_9^-$  with CN = 9,<sup>7,8</sup> and, finally, to the decacoordinated  $D_{10h}$  Ta $@B_{10}^-$  and  $D_{10h}$  Nb $@B_{10}^-$  with CN = 10 which has proven to be the highest coordination number in planar species observed to date.<sup>1,2</sup>

However, planar hypercoordination chemistry may go well beyond metal-centered monocyclic boron wheel clusters. The recent experimental characterization of perfect planar  $D_{9h}$  C<sub>18</sub> by atom manipulations and high-resolution atomic force microscopy<sup>9</sup> inspires us to coordinate transition metal centers with bare cyclo[*n*]carbon ring-like clusters ( $C_n$ ,  $n \ge 11$ ) to form metal-centered monocyclic carbon wheel complexes  $M \otimes C_n^{k+}$ . Very recently, the electronic properties of metal-carbon ring complexes such as Li@C<sub>18</sub> and MC<sub>16</sub> have been reported.<sup>10,11</sup> Carbon ( $[He]2s^22p^2$ ) is known to be effective ligand to various transition metals with a smaller covalent radius than boron ([He]2s<sup>2</sup>2p<sup>1</sup>), while group IIIB metals La, Y, and Sc with the valence electronic configurations of (n-1)d<sup>1</sup>ns<sup>2</sup> possess the largest covalent radii in the periodic table.12 Metal-centered monocyclic carbon wheel clusters  $M @C_n^{k+}$  (M = La, Y, Sc) are thus possible to have even higher coordination numbers than their boron counterparts  $M \otimes B_n^{k-}$ . In fact, two families of  $LaC_n^+$ monocations (n = 12-40) were observed in mobility measurements as early as in 1994, with a La atom inserted into the carbon ring for even-numbered clusters or attached to the inside or outside of the carbon ring for odd-numbered

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clusters.13 More detailed mass spectra were reported late on  $LaC_n^+$  in which  $LaC_{13}^+$  appeared to be a prominent species with the La center most likely attached inside a C<sub>13</sub> ring (ring Ib), while  $LaC_{14}^{+}$  was the only  $LaC_{2n}^{+}$  cluster with even number of C atoms for which ring Ib structure had a higher mass intensity than its competing isomer with a La atom inserted into a C<sub>14</sub> ring (ring Ia).14 Similar prominent mass peaks were also observed for YC13+, CeC13+, and ScC13+.15 Early density functional theory (DFT) calculations indicated that the most stable isomer of LaC<sub>13</sub><sup>+</sup> possessed a nearly-cumulenic structure with the La atom located at the center of the carbon ring.16,17 However, the unique role these experimentally observed  $LaC_n^+$ species (n = 13, 14) play in planar hypercoordination chemistry has largely omitted in previous investigations and their accurate geometrical and electronic structures and detailed La-C coordination bonding patterns remain to be fully evaluated using the state-of-the-art theoretical approaches to interpret their behaviors observed in experiments.

Detailed first-principles theory calculations performed in this work indicate that the experimentally observed La©C<sub>13</sub><sup>+</sup> (1) with a perfect  $D_{13h}$  symmetry and slightly off-centered La©C<sub>14</sub><sup>+</sup> (4) with a  $C_{2v}$  geometry achieve the record coordination numbers of CN = 13 and 11 in planar species reported to date. The enhanced stability of La©C<sub>13</sub><sup>+</sup> (1) originates from its  $\sigma + \pi$ dual aromaticity with 14 delocalized  $\sigma$  electrons and 14 delocalized  $\pi$  electrons (14  $\sigma$  + 14  $\pi$ ) each matching the 4*N* + 2 aromatic rule ( $N_{\sigma} = N_{\pi} = 3$ ).  $\sigma + \pi$  dually aromatic  $D_{13h}$  Ca©C<sub>13</sub> (2),  $C_{13v}$  Ac©C<sub>13</sub><sup>+</sup> (3),  $C_{2v}$  Y©B<sub>6</sub>C<sub>6</sub><sup>+</sup> (5), and  $C_{2v}$  Sc©B<sub>5</sub>C<sub>6</sub> (6) have also been obtained at first-principles theory level with CN = 13, 13, 12, and 11, respectively. The highly stable La©C<sub>13</sub><sup>+</sup> (1) is found to behave like a super-hydrogen monocation (H<sup>+</sup>) in its substituted complex compounds.

#### 2. Theoretical procedure

Extensive global-minimum (GM) searches were performed on  $LaC_{13}^{+}$ ,  $LaC_{14}^{+}$ ,  $CaC_{13}$ ,  $AcC_{13}^{+}$ ,  $YB_6C_6^{+}$ ,  $ScB_5C_6$ , and  $CeC_{13}^{+/2+}$ using the TGmin2 code18 at DFT level based on the constraint basin-hopping algorithm.<sup>19</sup> In total, more than 1000 stationary points with different spin multiplicities were probed for each species at PBE/DZVP level using the CP2K program.<sup>20,21</sup> Lowlying isomers were then fully optimized at the M06-2X and PBE0 level,<sup>22,23</sup> with the 6-311+G(d,p) basis sets for C, B, N, Ca, and Sc<sup>24</sup> and Stuttgart relativistic small-core pseudopotentials for La, Y, Ce, and Ac,<sup>25,26</sup> using the Gaussian-16 program suite.<sup>27</sup> Frequency analyses were performed to make sure all the optimized structures are true minima of the systems. Relative energies for the five lowest-lying isomers were further refined using the more accurate coupled cluster method with triple excitations CCSD(T)<sup>28-30</sup> implemented in Molpro<sup>31</sup> with the basis set of cc-pVTZ for C, B, N, Ca, and Sc and the Stuttgart smallcore pseudopotential for La, Y, and Ac. The optimized GM structures are summarized in Fig. 1 and more alternative isomers tabulated in Fig. S1-S6.<sup>†</sup> Natural bonding orbital (NBO) analyses were performed using the NBO 6.0 program.<sup>32</sup> Born-Oppenheimer molecular dynamics (BOMD) simulations were performed on  $LaC_{13}^{+}(1)$  for 30 ps using the CP2K software suite



**Fig. 1** Top and side views of the optimized La $@C_{13}^+$  (1), Ca $@C_{13}$  (2), Ac $@C_{13}^+$  (3), La $@C_{14}^+$  (4), Y $@B_6C_6^+$  (5), and Sc $@B_5C_6$  (6) at M06-2X level, with bond lengths indicated in Å.

at 300, 800, and 1000 K.<sup>21</sup> The anisotropy of the current-induced density (ACID)<sup>33</sup> analyses were performed using the ACID code, with the ring-current maps generated using POV-Ray 3.7.<sup>34</sup> The iso-chemical shielding surfaces (ICSSs)<sup>35,36</sup> were generated with the Multiwfn 3.8 code.<sup>37</sup> The UV-vis spectra were simulated using the time-dependent TD-DFT-M06-2X approach.<sup>38</sup>

#### 3. Results and discussion

We start from  $LaC_{13}^{+}$ , the most concerned species observed in  $LaC_n^+$  series (n = 12-40).<sup>13</sup> Encouragingly and interestingly, as shown in Fig. 1 and S1,† extensive GM searches indicate that  $LaC_{13}^{+}$  possesses the well-defined perfect planar GM of  $D_{13h}$  $La@C_{13}^{+}$  (1) which contains a La atom located exactly at the center of the C13 wheel ligand, with the optimized La-C coordination bond lengths of  $r_{\text{La-C}} = 2.70$  Å which are slightly longer than the sum of the self-consistent single-bond covalent radii of La and C (2.55 Å)<sup>12</sup> and C–C bond lengths of  $r_{C-C} = 1.29$  Å which lie between C=C double-bond (1.34 Å) and C≡C triple bond (1.20 Å), setting up the highest coordination number of CN = 13 in planar species reported to date.  $La@C_{13}^+$  (1) possesses the huge HOMO–LUMO gap of  $\Delta E_{gap} = 5.33$  eV at M06-2X level, well underlying its high chemical stability. The slightly distorted triplet  $C_s \operatorname{LaC_{13}}^+$  (<sup>3</sup>A') appears to be the second lowest-lying isomer lying 2.41 eV above the GM at CCSD(T). The seventh  $C_{2v}$  isomer with a La inserted into the  $C_{13}$  ring and the fifth  $C_{2v}$ isomer with a La attached to the outside of the  $C_{\rm 13}$  ring are found to lie 3.36 eV and 3.09 eV higher in energy than the GM at M06-2X, respectively. Extensive BOMD simulations indicate that  $La \otimes C_{13}^+$  (1) is highly dynamically stable at both 800 K and 1000 K, with the small average root-mean-square-deviations of RMSD = 0.11 and 0.13 Å and maximum bond length deviations of MAXD = 0.31 and 0.36 Å, respectively (Fig. S7<sup>†</sup>). No high-lying isomers were observed during the dynamical simulations.

Replacing the La center in  $La@C_{13}^+$  (1) with a Ca atom generates the charge-transfer neutral complex  $D_{13h}$  Ca@C<sub>13</sub> (2). The Ca center in Ca@C<sub>13</sub> (2) matches the C<sub>13</sub> wheel ligand

perfectly both electronically and geometrically though it has a much lower Wiberg bond index of  $WBI_{Ca} = 0.26$  than La in  $La@C_{13}^{+}(1)$  where  $WBI_{La} = 1.76$  (Table 1). The Ca–C coordination interaction with the low bond order of  $WBI_{Ca-C} = 0.02$  in  $Ca^{2+} @C_{13}^{2-}$  (2) is thus almost purely ionic. Using an Ac atom which has a larger atomic radius than La to replace La in La $\mathbb{O}C_{13}^{+}(1)$ , the slightly buckled  $C_{13v}$  Ac $\mathbb{O}C_{13}^{+}(3)$  is generated in which the Ac atom lies 0.59 Å above the  $C_{13}$  ring, with the periphery C–C distances of  $r_{C-C} = 1.29$  Å remaining basically unchanged. The experimentally observed prominent  $ScC_{13}^{+}$  and  $YC_{13}^{+}$  monocations<sup>15</sup> have severely off-centered  $C_{2v}$  Sc $OC_{13}^{+}$  $({}^{1}A_{1})$  (with CN = 8) and  $C_{2v}$  Y©C<sub>13</sub><sup>+</sup>  $({}^{1}A_{1})$  (with CN = 9) GM structures due to strong ring strains, respectively, while the open-shell  $\operatorname{CeC}_{13}^+$  possesses a slightly distorted GM  $C_{2v} \operatorname{Ce} \mathbb{O} \operatorname{C}_{13}^+$  $({}^{2}B_{2})$  which has practically a  $D_{13h}$  symmetry (Fig. S8<sup>†</sup>). With one more valence electron detached, the Ce©C<sub>13</sub><sup>2+</sup> dication isovalent with La $@C_{13}^+$  (1) has indeed a perfect  $D_{13h}$  GM (Fig. S8<sup>†</sup>).

It is natural to ask at current stage whether it is possible to form metal-centered monocyclic carbon wheel clusters with coordination numbers greater than thirteen (*i.e.*, CN > 13). We carefully checked the hypercoordination chemistry of the experimentally observed La©C<sub>14</sub><sup>+</sup> in this work. Although a bare C14 has a perfect C7h acetylenic structure,39 with a La atom added in, the La-doped LaC<sub>14</sub><sup>+</sup> possesses the off-centered planar GM of  $C_{2v}$  La©C<sub>14</sub><sup>+</sup> (4) which has the actual coordination number of CN = 11 (Fig. 1 and S4<sup>†</sup>). The three C atoms on the top part of La $OC_{14}^+$  (4) with La–C distances great than 3.1 Å have the practically negligible La-C coordination bond orders (with WBI<sub>La-C</sub>  $\approx$  0.00). Using Ac, the largest actinide metal in the periodical table, to replace La<sup>12</sup> an elongated planar  $D_{2h}$ Ac $\mathbb{O}C_{14}^+$  (Fig. S8<sup>†</sup>) with CN = 12 is generated. A  $C_{14}$  ring is obviously too big in size to host a transition-metal atom at its geometrical center comfortably to form a perfect wheel complex  $D_{nh}$  M©C<sub>n</sub> with the same M–C coordination bonding distances. We conclude that CN = 13 is the highest coordination number in metal-centered monocyclic carbon wheel clusters  $M \otimes C_n^+$ with effective M-C coordination interactions.

Introducing certain numbers of B atoms into the carbon rings generates more structural diversities in B–C binary wheel ligands  $B_mC_n$  with CN  $\leq$  13. As examples, the metal-centered

monocyclic C–B binary wheel complexes  $Y @B_6C_6^+$  (5) and  $Sc @B_5C_6$  (6) in Fig. 1 and  $C_{2v} La @BC_{12}$  in Fig. S8† as the GMs of the systems possess the coordination numbers of CN = 12, 11, and 13, respectively. These planar C–B binary wheel complexes have larger C–B and B–B periphery distances than the corresponding C–C distances in  $La @C_{13}^+$  (1) because B has a larger covalent radius than C.

To interpret the high stabilities of these hypercoordinated planar species, we performed detailed NBO and molecular orbital analyses on  $\text{La}^{\odot}\text{C}_{13}^{+}(1)$ ,  $\text{Y}^{\odot}\text{B}_{6}\text{C}_{6}^{+}(5)$ , and  $\text{Sc}^{\odot}\text{B}_{5}\text{C}_{6}(6)$ . As tabulated in Table 1, the La center in  $\text{La}^{\odot}\text{C}_{13}^{+}(1)$  possesses the natural atomic charge of  $q_{\text{La}} = + 2.08$  |e|, electronic configuration of  $\text{La}[\text{Xe}]4\text{f}^{0.43}5\text{d}^{0.41}\text{6s}^{0.02}$ , and total Wiberg bond order of WBI<sub>La</sub> = 1.76. The C atoms on the C<sub>13</sub> wheel ligand have the total Wiberg bond orders of WBI<sub>C</sub> = 3.99 and C–C bond orders of WBI<sub>C-C</sub> = 1.73, revealing the cumulenic nature of the complex. Obviously, the La center donates its 6s electron almost completely to the C<sub>13</sub> ligand. The La–C<sub>13</sub> coordination



Fig. 2 Delocalized  $\pi$ - and  $\sigma$ -CMOs of (a) La $\otimes$ C<sub>13</sub><sup>+</sup> (1), (b) Y $\otimes$ B<sub>6</sub>C<sub>6</sub><sup>+</sup> (5), and (c) Sc $\otimes$ B<sub>5</sub>C<sub>6</sub> (6), in comparison with the delocalized  $\pi$ -CMOs of (d)  $D_{6h}$  C<sub>6</sub>H<sub>6</sub>.

Table 1 Calculated HOMO-LUMO gaps $\Delta E_{qap}/eV$ , natural atomic charges $q_M$ , electronic configurations, and Wiberg bond indexes WBI <sub>M</sub> of the
metal centers M and Wiberg bond indexes of the $M-C$ ( $WBI_{M-C}$ ) and $M-B$ ( $WBI_{M-B}$ ) coordination interactions and NICS values 1.0 Å above the
metal centers (NICS (1)) of the $M^{(x)}$ $\otimes B_m C_n^{k\pm}$ complex series (1–6) at M06-2X level

$\mathbf{M}^{(x)} \odot \mathbf{B}_m \mathbf{C}_n^{k\pm}$	CN	$\Delta E_{ m gap}/ m eV$	$q_{\mathbf{M}}$	Electronic configurations of central metal M	WBI <sub>M</sub>	WBI <sub>M-C</sub>	WBI <sub>M-B</sub>	NICS (1)
$D_{13h} \text{ La} \odot C_{13}^{+}(1)$	13	5.33	2.08	$[Xe] 4f^{0.43}5d^{0.41}6s^{0.02}$	1.76	0.14	_	-58.54
$D_{13h}$ Ca©C <sub>13</sub> (2)	13	5.63	1.87	[Ar] 3d <sup>0.09</sup> 4s <sup>0.03</sup> 4p <sup>0.01</sup>	0.26	0.02	_	-44.68
$C_{13v} \operatorname{Ac} \mathbb{O} C_{13}^{+} (3)$	13	5.26	2.38	[Rn] 5f <sup>0.19</sup> 6d <sup>0.35</sup> 7s <sup>0.03</sup>	1.22	0.10	_	-55.16
$C_{2v}$ La©C <sub>14</sub> <sup>+</sup> (4)	11	3.72	2.15	$[Xe] 4f^{0.28}5d^{0.50}6s^{0.03}$	1.64	0.07-	_	-29.21
				2 3		0.16		
$C_{2v} \operatorname{Y} \mathbb{O} \operatorname{B_6 C_6}^+(5)$	12	5.70	2.12	$[Kr] 5s^{0.06}4d^{0.70}5p^{0.01}$	1.67	0.10-	0.09-	-24.08
						0.22	0.12	
$C_{2v}$ Sc©B <sub>5</sub> C <sub>6</sub> (6)	11	5.32	1.94	$[Ar] 3d^{0.87}4s^{0.07}4p^{0.02}$	1.94	0.13-	0.13-	-29.62
						0.29	0.14	

interactions mainly originate from contributions involving the 5d and 4f atomic orbitals of the La center, as indicated in the degenerated  $\pi$ -HOMO/HOMO',  $\pi$ -HOMO-2/HOMO-2',  $\sigma$ -HOMO-1/HOMO-1', and  $\sigma$ -HOMO-3/HOMO-3 in Fig. 2(a). Although each La–C coordination interaction in La $^{\circ}$ C<sub>13</sub><sup>+</sup> (1) has a relatively low bond order (WBI<sub>La–C</sub> = 0.14), the thirteen equivalent La–C coordination bonds function together to effectively stabilize the hypercoordinated cluster, making it the well-defined GM of the system observed solely in gas-phase experiments.<sup>13,14</sup> Similar situations happen in Ac $^{\circ}$ C<sub>13</sub><sup>+</sup> (3), Y $^{\circ}$ B<sub>6</sub>C<sub>6</sub><sup>+</sup> (5), and Sc $^{\circ}$ B<sub>5</sub>C<sub>6</sub> (6).

The 14 totally delocalized canonical molecular orbitals (CMOs) of La $^{\circ}$ C<sub>13</sub><sup>+</sup> (1) are collectively shown in Fig. 2(a), including 7 delocalized  $\pi$ -CMOs and 7 delocalized  $\sigma$ -CMOs. Its remaining 13 σ-CMOs correspond to 13 two-center-two-electron (2c-2e) C–C  $\sigma$  bonds along the periphery of the C<sub>13</sub> ligand (Fig. S9<sup>†</sup>). The degenerate f-type HOMO/HOMO', d-type HOMO-2/HOMO-2' and p-type HOMO-5/HOMO-5' and non-degenerate s-type HOMO-6 possess three, two, one, and zero nodal surfaces, respectively, forming an out-of-plane 14  $\pi$  electron system matching the  $4N_{\pi}$  + 2 aromatic rule with  $N_{\pi}$  = 3. Similarly, the degenerate HOMO-1/HOMO-1', HOMO-3/HOMO-3', and HOMO-4/HOMO-4' and non-degenerate HOMO-7 form an in-plane 14  $\sigma$  electron system matching the  $4N_{\sigma}$  + 2 aromatic rule with  $N_{\sigma} = 3$ . Such a unique 14  $\sigma$  + 14  $\pi$  electronic configuration renders  $\sigma + \pi$  dual aromaticity to La©C<sub>13</sub><sup>+</sup> (1), effectively stabilizing the monocation observed in experiments, similar to the situation in the  $\sigma + \pi$  dually aromatic  $D_{10h}$  Ta©B<sub>10</sub><sup>-</sup> which possesses a 10  $\sigma$  + 6  $\pi$  electronic configuration.<sup>1</sup> It is noticed that metal-centered wheel  $D_{13h}$  La $\odot C_{13}^+$  (1) has the same numbers of delocalized  $\sigma$  and  $\pi$  electrons (14  $\sigma$  + 14  $\pi)$  as the highly stable ring-like acetylenic C<sub>7h</sub> C<sub>14</sub>.<sup>39</sup>

As shown in Fig. 2(b) and (c), both the metal-centered monocyclic C–B binary wheel clusters  $Y @B_6 C_6^+$  (5) and  $Sc @B_5 C_6$  (6) have 5 delocalized  $\pi$ -CMOs and 5 delocalized  $\sigma$ -CMOs, forming an out-of-plane 10  $\pi$  electron system conforming to the  $4N_{\pi} + 2 \pi$ -aromatic rule ( $N_{\pi} = 2$ ) and an in-plane 10  $\sigma$  electron system conforming to  $4N_{\sigma} + 2 \sigma$  aromatic rule ( $N_{\sigma} = 2$ ). Such 10  $\sigma + 10 \pi$  electronic configurations make both Y@B<sub>6</sub>C<sub>6</sub><sup>+</sup> (5) and Sc@B<sub>5</sub>C<sub>6</sub> (6)  $\sigma + \pi$  dually aromatic in nature, similar to but with more delocalized electrons than the previously reported Co@B<sub>8</sub><sup>-</sup> and Ru@B<sub>9</sub><sup>-</sup> with 6  $\sigma + 6 \pi$  delocalized electrons and Ta@B<sub>10</sub><sup>-</sup> and Nb@B<sub>10</sub><sup>-</sup> with 10  $\sigma + 6 \pi$  delocalized electrons.

The La–C<sub>13</sub> coordination interactions in La©C<sub>13</sub><sup>+</sup> (1) mainly originate from d–p and f–p  $\sigma$ -coordination interactions involving the in-plane La  $5d_{x^2-y^2}/5d_{xy}$  and  $4f_{x(x^2-3y^2)}/4f_{y(y^2-3x^2)}$ atomic orbitals and C  $2p_x/2p_y$  hybridized atomic orbitals, as demonstrated in the degenerated HOMO-3/HOMO-3' (5d–2p  $\sigma$ coordination) and HOMO-1/HOMO-1' (4f–2p  $\sigma$ -coordination) in Fig. 2(a), respectively. Quantitatively, the La center contributes more to the delocalized in-plane  $\sigma$ -CMOs (2.81–7.63%) than to the delocalized out-of-plane  $\pi$ -CMOs (0.08–3.66%). The delocalized in-plane  $\sigma$ -CMOs thus dominate the La–C coordination bonding interactions. Similarly, the in-plane 4d–2p  $\sigma$ -coordination interactions (HOMO-2 and HOMO-3) in Y©B<sub>6</sub>C<sub>6</sub><sup>+</sup> (5) and 3d–2p  $\sigma$ -coordination interactions (HOMO-2 and HOMO-3) in



Fig. 3 (a) Calculated iso-chemical shielding surfaces (ICSSs) of  $La@C_{13}^+$  (1),  $Y@B_6C_6^+$  (5), and  $Sc@B_5C_6$  (6), compared with that of aromatic benzene  $C_6H_6$ . Yellow and green regions stand for chemical shielding and de-shielding areas, respectively. (b) Calculated  $\pi$ - and  $\sigma$ -ring current maps of  $La@C_{13}^+$  (1),  $Y@B_6C_6^+$  (5), and  $Sc@B_5C_6$  (6), respectively, in comparison with the  $\pi$ -ring current map of  $D_{6h}C_6H_6$ . The external magnetic field is perpendicular to the wheel plane. The red arrows represent directions and magnitudes of the ring currents at various positions on the ACID iso-surfaces.

Sc©B<sub>5</sub>C<sub>6</sub> (6) dominate the metal-ligand interactions in the two 10  $\sigma$  + 10  $\pi$  systems.

The aromatic nature of  $La@C_{13}^+$  (1),  $Y@B_6C_6^+$  (5), and  $Sc@B_5C_6$  (6) is further evidenced by their calculated nucleusindependent chemical shift (NICS) values. Based on the calculated NICS-ZZ components, Fig. 3(a) depicts the ICSS surfaces of 1, 5, and 6 with the Z-axis perpendicular to the molecular planes to illuminate the chemical shielding around the metal centers, in comparison with that of the prototypical aromatic benzene  $(C_6H_6)$ . Obviously, the space inside the  $C_{13}$ ,  $B_6C_6$ , or  $B_5C_6$  rings in horizontal direction or within about 1.0 Å above the metal centers in vertical direction belong to chemical shielding regions with negative NICS-ZZ values (highlighted in yellow), while the chemical de-shielding areas with positive NICS values (highlighted in green) are located outside the wheel ligands in horizontal direction. Fig. 3(a) clearly shows the ICSS surfaces of these metal-centered planar complexes are exactly similar to that of the aromatic benzene.

Calculating the anisotropy of current-induced density (ACID)<sup>34</sup> is an effective approach to graphically display the ring currents induced by an external magnetic field in vertical directions perpendicular to the molecular planes. ACID can be decomposed into  $\sigma$  and  $\pi$  components separately. Fig. 3(b) clearly indicates the  $\pi$ -ring current maps of La $\mathbb{O}C_{13}^+$  (1),  $Y \otimes B_6 C_6^+$  (5), and  $S \otimes C_6 (6)$  are extremely similar to the corresponding  $\pi$ -ring current map of  $\pi$ -aromatic benzene C<sub>6</sub>H<sub>6</sub>. Besides, in contrast to benzene which possesses no delocalized  $\sigma$ -electrons, La©C<sub>13</sub><sup>+</sup> (1), Y©B<sub>6</sub>C<sub>6</sub><sup>+</sup> (5), and Sc©B<sub>5</sub>C<sub>6</sub> (6) also exhibit strong  $\sigma$ -ring currents, rendering additional  $\sigma$ -aromaticity to stabilize the systems, as shown in Fig. 3(b) (see Fig. S10<sup>+</sup> for high-resolution  $\pi$ - and  $\sigma$ -ring current maps of **1**, **5**, and **6**). The observation of both  $\sigma$ - and  $\pi$ -diatropic ring currents in 1, 5, and **6** well supports the  $\sigma + \pi$  dually aromatic nature of these planar hypercoordinated complexes.

Based on the planar hypercoordinated species discussed above, we develop a universal electronic design principle for  $\sigma$  +  $\pi$  dually aromatic metal-centered monocyclic boron, carbon, or boron-carbon binary wheel clusters  $\mathbf{M}^{(x)} \odot \mathbf{B}_m \mathbf{C}_n^{\ k\pm}$ . There exist m+ n 2c-2e  $\sigma$  bonds along the periphery of the monocyclic  $\mathbf{B}_m \mathbf{C}_n$ wheel ligand. With each C atom providing two delocalized electrons and each B atom contributing one delocalized electron, the total number of the delocalized electrons in  $\mathbf{M}^{(x)} \odot \mathbf{B}_m \mathbf{C}_n^{\ k\pm}$  matches the requirement of  $\sigma + \pi$  dual aromaticity of the system:

$$m + 2n + x \pm k = L \ (L = 12, 16, 20, \text{ or } 28)$$
 (1)

where *L* is the total number of delocalized  $\sigma$  and  $\pi$  electrons each matching the 4N + 2 rule and x stands for the formal valence of the central atom M. Such an electronic design principle covers the previously observed  $D_{7h} B @B_7^{2-}$ ,  $D_{8h} B @B_8^{-}$ ,  $D_{9h}$  $Co@B_8^-$ , and  $D_{9h} Ru@B_9^-$  where L = 12 and x = 3, 3, 3, and 2and  $D_{10h}$  Ta©B<sub>10</sub><sup>-</sup> and  $D_{10h}$  Nb©B<sub>10</sub><sup>-</sup> where L = 16 and x = 5and 5, respectively. For the currently discussed  $La@C_{13}^{+}$  (1), Ca $^{\odot}C_{13}(2)$ , and Ac $^{\odot}C_{13}^{+}(3)$ , L = 28 and x = 3, 2, and 3, while for  $Y \otimes B_6 C_6^+$  (5) and  $S \otimes B_5 C_6$  (6), L = 20 and x = 3 and 3, respectively. The slightly buckled  $C_{13v}$  Ac $\mathbb{C}C_{13}^+$  (3) follows the same electronic design principle though the Ac center in it slightly mismatches the  $C_{13}$  wheel ligand in geometry. Eqn (1) can be easily extended to  $\sigma + \pi$  dually aromatic B–C–N ternary wheel complexes  $M^{(x)} \odot B_m C_n N_l^{k\pm}$  where it reads:  $m + 2n + 3l + x \pm k =$ 28, as demonstrated in the cases of  $C_{\rm s}$  La©BNC<sub>11</sub><sup>+</sup> and  $C_{\rm s}$  $La@B_4N_4C_5^+$  (Fig. S8<sup>†</sup>) in which each N atom contributes three valence electrons to the delocalized systems. However, extensive test calculations indicate that inclusion B or N atoms in the wheel ligands does not help to improve the maximum coordination numbers of the systems.

The highly stable  $La@C_{13}^+$  (1) with a wide HOMO-LUMO gap and fully occupied bonding inner-shell CMOs can be used as a super-hydrogen monocation (H<sup>+</sup>) to form various substituted multi-nuclei complexes. Typical examples include  $C_{13v}$  [La@C<sub>13</sub>] X (X = F, Cl, Br) (7),  $D_{13h}$  [La@C<sub>13</sub>]<sup>+</sup>L<sub>2</sub> (L = Ar, Kr) (8),  $C_{2v}$ [La@C<sub>13</sub>]<sub>2</sub>O (9), and  $C_{3v}$  N[La@C<sub>13</sub>]<sub>3</sub> (10) (Fig. 4) which can be derived from the parent species  $C_{\infty v}$  HX,  $D_{\infty h}$  H<sup>+</sup>L<sub>2</sub>,  $C_{2v}$  H<sub>2</sub>O, and  $C_{3v}$  NH<sub>3</sub> by substituting H<sup>+</sup> monocation(s) with La@C<sub>13</sub><sup>+</sup> (1) unit(s), respectively, presenting the viable possibility to form complex compounds with multiple hypercoordinated metal centers.

Finally, as examples, the simulated IR, Raman, and UV-vis spectra of  $D_{13h}$  La©C<sub>13</sub><sup>+</sup> (**1**) and  $C_{2v}$  Y©B<sub>6</sub>C<sub>6</sub><sup>+</sup> (5) are presented in Fig. S11† to facilitate their spectroscopic characterizations. La©C<sub>13</sub><sup>+</sup> (**1**) possesses highly simplified spectra due to its perfect  $D_{13h}$  symmetry. It has two main IR peaks at 39 ( $a_2^{"}$ ) and



Fig. 4 Optimized structures of  $[La \odot C_{13}]X$  (7) (X = F, Cl, Br),  $[La \odot C_{13}]^+L_2$  (8) (L = Ar, Kr),  $[La \odot C_{13}]_2O$  (9), and N $[La \odot C_{13}]_3$  (10) at M06-2X level.

866  $(e'_1) \text{ cm}^{-1}$ , three Raman peaks at 272  $(e'_2)$ , 664  $(a'_1)$ , and 1280  $(e'_2) \text{ cm}^{-1}$ , and three UV absorption peaks at 156  $(A'_1)$ , 174  $(E'_2)$ , and 200  $(E'_2)$  nm, respectively.  $C_{2v} \text{ Y} \otimes B_6 \text{C}_6^+$  (5) has more complicated spectra, with two main IR absorption peaks at 499  $(b_1)$  and 588  $(b_2) \text{ cm}^{-1}$ , two major Raman scattering bands at 211  $(b_2)$  and 623  $(a_1) \text{ cm}^{-1}$ , and five UV absorption bands around 134  $(^1B_2)$ , 161  $(^1A_1)$ , 191  $(^1B_2)$ , 207  $(^1A_1)$ , and 219  $(^1B_2)$  nm, respectively.

#### 4. Conclusions

Extensive first-principles theory calculations performed in this work unveil the highest coordination numbers of CN = 13 in  $La@C_{13}^{+}(1)$ ,  $Ca@C_{13}(2)$ , and  $Ac@C_{13}^{+}(3)$ , CN = 12 in  $Y@B_6C_6^{+}$ (5), and CN = 11 in La $@C_{14}^+$  (4) and Sc $@B_5C_6$  (6) reported to date in planar species, effectively enriching the structural and bonding patterns in planar hypercoordination chemistry.  $\sigma + \pi$ dually aromatic metal-centered monocyclic B-C binary wheel complexes  $M^{(x)} \odot B_m C_n^{k\pm}$  follow the electronic design principle of  $m + 2n + x \pm k = L$  (L = 12, 16, 20, or 28) which can be readily extended to more complicated systems containing other periphery atoms rather than B and C. The recent experimental identification of the sp-hybridized molecular carbon allotrope,  $\pi$ -aromatic C<sub>18</sub>, induced by atom manipulations,<sup>9</sup> presents the possibility to synthesize and characterize the  $\sigma$  +  $\pi$  dually aromatic  $La@C_{13}^+(1)$  and its complex compounds to open a new area in planar hypercoordination chemistry, catalysis, and materials science.

## Conflicts of interest

There are no conflicts to declare.

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