A bifunctional strategy towards experimentally (synthetically) attainable molecules with planar tetracoordinate carbons

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Bifunctional strategy, including efficiently utilizing valence electrons and offering steric protection, has been proposed to advance C2Al4 global minimum with double planar tetracoordinate carbons (ptC) to a new family of ptC molecules which could be promising for synthetic realization.

Stabilizing the extremely unfavorable planar tetracoordinate carbon (ptC) has always been a challenge.1,2 The first prediction of the ptC molecule (1,1-dilithiocyclopropane)3 encouraged strategic developments, and more computational and experimental realizations.4–9 In the past decade, the great efforts from computational chemists have advanced the ptC chemistry to the general planar hypercoordinate carbons (phC) where the coordination numbers of the flat carbons are more than three (hence, including ptC). Numerous computed phC molecules have emerged.4–17 In contrast, experimentally (synthetically, in particular) accessible phC molecules are rare. Only several transition metal and lithium-containing phC compounds have been synthesized.4 The recently reported CAl42 with its analogues were only observed in the gas phase.18–20 In these computed phC molecules, the electron deficient elements (e.g., B, Al, Li, and transition metals) were frequently used as electron acceptors and the ligands were mostly exposed externally. Exposing such electron deficient atoms results in their high reactivity, and the lack of “anchors” to attach protecting groups in these species further limits the operations of conventional synthetic strategies. In many cases the computed phC molecules are not the global minima, which is a concern; this also presents difficulties for experimental realizations even in the gas phase. For example, the global minimum, CAl42– and its analogues were only observed in the gas phase.18–20 In these computed phC molecules, the electron deficient elements (e.g., B, Al, Li, and transition metals) were frequently used as electron acceptors and the ligands were mostly exposed externally. Exposing such electron deficient atoms results in their high reactivity, and the lack of “anchors” to attach protecting groups in these species further limits the operations of conventional synthetic strategies. In many cases the computed phC molecules are not the global minima, which is a concern; this also presents difficulties for experimental realizations even in thegas phase. For example, the global minimum, CAl42– and its analogues have been observed, but the efforts to detect the hypercoordinated CB7– and CB62– (not global minima) led to the more favorable species in which the carbons prefer locating on the edges or vertexes.21,22 Therefore, the more challenging task faced by computational chemists is to design phC species attainable experimentally or synthetically in particular (i.e., not just be observable in the gas phase). This will help develop the potential applications of such exotic carbon bonding patterns. Here we propose a bifunctional strategy to advance our newly uncovered global minimum species (D2h, C2Al4) with double planar tetracoordinate carbons (dptCs) to those which are promising for synthetic realizations.

Using Gaussian 03,23 most of the reported structures were optimized at the B3LYP/6-311++G** level, while the largest molecule at the B3LYP/6-31G* level. Subsequent frequency analyses were performed to verify the optimized structures to be minima and the unscaled harmonic frequency values were used to evaluate their relative thermodynamic stabilities. The smallest species, C2Al4H8 (1, Fig. 1) was reoptimized at the B3LYP/aug-cc-pVTZ and MP2/aug-cc-pVTZ levels to calibrate suitability of the basis sets and the B3LYP method. The electronic structures were analyzed using the NBO.5.0 program24 at the B3LYP level. The Born–Oppenheimer molecular dynamic (BOMD)25–27 simulations were run at B3LYP/6-31G* level.

Continuing our interest in phC chemistry, we recently got an intriguing finding, C2Al4 is the simplest neutral global minimum.28 Comparing the electronic structure of C2Al4 with

Fig. 1 Optimized structures of 1–7 at B3LYP/6-311++G** and 8 at B3LYP/6-31G*. The key bond lengths are given in Å. The numbers after the point groups are the smallest frequencies in cm–1 and the HOMO–LUMO gaps in eV. The values in the parentheses, and brackets given in 1, are at B3LYP/aug-cc-pVTZ and MP2/aug-cc-pVTZ, respectively. Color code, yellow: Al, cyan: F, red: O, blue: N, grey: C, pink: Be, and white: H.

abcdefg
those of \( \text{CAI}_{2}^{2-} \) and \( \text{CAI}_{4}^{+} \) led to a succinct equivalence relationship: the central CC moiety in \( \text{C}_2\text{Al}_4 \) is equivalent to the carbon centers in \( \text{CAI}_{2}^{2-} \) and \( \text{CAI}_{4}^{+} \). \( \text{C}_2\text{Al}_4 \) also contrasts to ethylene by possessing two different molecular orbitals (MO’s). The two MO’s of ethylene, related to the C–H and C–C σ bonds are replaced by the peripheral Al4 and four-center-involved (CAI\(_{BR}\)/CAI\(_{BR}\)) bonding MO’s in \( \text{C}_2\text{Al}_4 \). Hereafter, Al\(_{BR}\) and Al\(_{TE}\) represent the bridging and terminal Al atoms, respectively. Although \( \text{C}_2\text{Al}_4 \) is the global minimum, we speculated that such species can only be detected as transient species in the gas phase and is probably difficult to attain by conventional synthesis due to the reasons mentioned above.

\( \text{C}_2\text{Al}_4 \) is a global minima, but it does not have optimal electronic structure: the eight more valence electrons than ethylene occupy four non-bonding MO’s and are not effectively utilized for chemical bonding. When designing boraplanes and their analogues, Wang and Schleyer proposed to direct the two non-bonding electrons on ptC into a peripheral B4 bonding orbital in boraplanes (or C–B bonds in their analogs) to utilize the otherwise “wasted” electrons and thus to achieve perfect ptC arrangements.29,30 Following this idea, we expect that directing the eight non-bonding electrons in \( \text{C}_2\text{Al}_4 \) into bonding MO’s will benefit its stability and thus improve its chemical accessibility. Moreover, applying the equivalence relationship to our31 recently designed \( \text{CBe}_4\text{H}_4 \) also led to a \( D_{2h} \text{C}_2\text{Be}_4\text{H}_4 \) minimum (2). Hence it is anticipated that further replacement of BeH groups in 2 with the valence isoelectronic AlH2 will result in a true \( D_{2h} \) minimum with perfect dptCs. 1. This expectation was confirmed by computations at the B3LYP/6-311++G** , B3LYP/aug-cc-pVTZ and MP2/aug-cc-pVTZ levels. The key geometric parameters at these three levels are compared in Fig. 1, which indicates that the B3LYP/6-311++G** is an appropriate level in describing this system. Thus, computational results at this level (B3LYP/6-311++G**) are discussed hereafter, unless otherwise specified.

The four occupied non-bonding MO’s (column 2 in Fig. 2) of the isolated \( \text{C}_2\text{Al}_4 \) correlate to the eight occupied MO’s (columns 3 and 4) of 1. Although one can observe somewhat non-bonding characters in the eight MO’s of 1, it is apparent that the electrons are much more efficiently utilized for bonding. This is strongly supported by the heat formation of \( \text{C}_2\text{Al}_4 + \text{H}_2 \) to 1, \(-75.4 \text{ kcal mol}^{-1} \) (\( \Delta H \)). The free energy changes \( (\Delta G) \) are given in Table 1 for comparison. The characteristic occupied orbitals of \( \text{C}_2\text{Al}_4 \) (the first three MO’s in column 1) have their counterparts in column 5. The bonding interactions between C and Al\(_{TE}\) in the LUMO of \( \text{C}_2\text{Al}_4 \) indicate its great potential to improve its chemical accessibility. Moreover, applying the equivalence relationship to our31 recently designed \( \text{CBe}_4\text{H}_4 \) (analogs) to utilize the otherwise “wasted” electrons and thus the more efficient utilization of the valence electrons than in \( \text{C}_2\text{Al}_4 \). This is also unveiled by NBO electronic structure analyses. The Wiberg bond indices (WBI’s) of C–Al\(_{TE}\)/C–Al\(_{BR}\) bonds in \( \text{C}_2\text{Al}_4 \) are 0.53/0.62 (Table 2), while the values in 1 are increased to 0.65/0.67. The ionization potentials between the central CC moiety and the Al4 ring are also strengthened. Due to the relatively large electronegativity of H compared to Al, hydrogen atoms bear negative charges, \(-0.33 \text{ e/AlH} \) and \(-0.32 \text{ e/H–AlBR} \), resulting in much larger positive charges on Al\(_{TE}/\text{AlBR}\) (1.22e/1.14e) than the 0.68e/0.66e in \( \text{C}_2\text{Al}_4 \). Although the central carbon atoms in 1 bear less negative charges, \(-1.07 \text{ e} \) vs. \(-1.34 \text{ e} \) in \( \text{C}_2\text{Al}_4 \), the overall ionic interactions in 1 are increased on the basis of the Coulomb attraction consideration. Therefore the total interactions between Al and C atoms are enhanced, consistent with the more compact \( \text{C}_2\text{Al}_4 \) core in 1.

The electronic and geometric analyses only indicate the optimal electronic structure of 1. The thermodynamic stability is important for the experimental realization. Eqn (1)–(3) (Table 1), give good estimations on its thermodynamic stability. The full hydrogenation of \( \text{C}_2\text{Al}_4 \) to 1 (eqn (1)) is

\[
\begin{align*}
\text{C}_2\text{Al}_4 + 4 \text{H}_2 & \rightarrow 1 \quad (1) \quad -75.4\text{ kcal mol}^{-1} \quad -40.0\text{ kcal mol}^{-1} \quad -46.5\text{ kcal mol}^{-1} \\
\text{C}_2\text{H}_4 + \text{AlH}_4 & \rightarrow 1 \quad (2) \quad -65.2\text{ kcal mol}^{-1} \quad -35.9\text{ kcal mol}^{-1} \quad -56.6\text{ kcal mol}^{-1} \\
\text{C}_2\text{H}_2 + \text{AlH}_6 & \rightarrow 1 \quad (3) \quad -36.9\text{ kcal mol}^{-1} \quad -23.3\text{ kcal mol}^{-1} \quad -28.8\text{ kcal mol}^{-1}
\end{align*}
\]

\( ^{a} \text{B3LYP/6-311++G**} \). \( ^{b} \text{MP2/aug-cc-pVTZ} \).
exothermic by 75.4 kcal mol$^{-1}$ ($\Delta H$), and the free energy of eqn (1) is $-41.0$ kcal mol$^{-1}$. Note that C$_2$Al$_4$ was confirmed to be a global minimum.$^{29}$ More realistic estimations are given by eqn (2) and (3). Relative to the experimentally available C$_2$H$_4$ + AlH$_4$ and C$_2$H$_2$ + AlH$_6$, I is 46.3 and 31.7 kcal mol$^{-1}$ ($\Delta H$) more stable, respectively. In comparison with the MP2/aug-cc-pVTZ results, the B3LYP underestimates the favorability of I by several kcal mol$^{-1}$ (Table 1). The exothermicity indicates the good thermodynamic stability of I. The ethylene-like isomer (3, as shown in Fig. 1) of I has six imaginary frequencies and is 41.6 kcal mol$^{-1}$ more stable, respectively. In contrast to various experimentally realized species, we further examined its thermal stability by two sets of BOMD (Born–Oppenheimer molecular dynamics) simulations at 323 and 373 K, respectively, starting at I. As reflected by structural evaluation described by the RMSD (root-mean-square deviation relative to I) vs. simulation time (Fig. 3), I was well maintained in the 35 ps simulation. The dynamic simulations imply that there are no nearby low-lying isomers which can be visited by crossing low energy barriers, though they can not tell whether I is a global minimum. Therefore it can be safely concluded that I has adequate thermal and thermodynamic stability for experimental realizations.

Substituting Al’s in C$_2$Al$_4$ by AlH$_2$ groups is a bifunctional approach. On the one hand, the additional eight Al–H bonds can utilize the eight otherwise “wasted” electrons of C$_2$Al$_4$, but on the other hand, the AlH$_2$ groups provide “anchors” to attach protecting groups in the attempts to synthesize such molecules. The use of protecting groups is a common strategy to prepare reactive species such as bottleable carbene.$^{34,35}$ I can be served as a prototype for a series of dptC molecules. Elaborating I with proper substituents gives new dptC molecules. Replacing all the hydrogen atoms in I with CH$_3$, NH$_2$, OH and F groups lead to minima, 4(D$_{3h}$), 5(D$_{2h}$), 6(D$_2$), and 7(D$_{2h}$), respectively. Note that the modes corresponding to the smallest vibrational frequencies (values are shown in Fig. 1) are not related to the distortion motions of the planar C$_2$Al$_4$ cores. The C$_2$Al$_4$ cores in all the derivatives maintain structures with perfect dptCs (i.e. C$_2$Al$_4$ core lie in the same plane). Similar to I, the total WBI’s of CC–AlTE/CC–AlBR (the WBI between CC moiety and AlTE or AlBR) and NBO charges on C and AlTE/AlBR atoms (Table 2) reflect the enhanced covalent and ionic bonding interactions in these derivatives, with respect to the isolated C$_2$Al$_4$. The WBI’s and NBO charges also indicate that, as the electronegativity of the central atoms of the substituents increase (C to N, to O, and to F), the covalent bonding interactions decrease and the ionic interactions increase (see Table 2). The overall interactions between the CC moiety and the Al$_4$ ring, as indicated by the shortened C–AlTE/CC–AlBR bonds, increase as the central atom of the substituent moves from C to F in the periodical table.

One can also attach bulky protecting groups to stabilize these species or hinder their possible high reactivity. Molecule 8 in Fig. 1 is the B3LYP/6-31G* optimized structure with the t-Bu groups. As expected, the C$_2$Al$_4$ core in 8 expands slightly due to the crowded substituents (the C–AlTE/CC–AlBR bond lengths, 1.960/2.111(2.096) Å vs. 1.913 and 2.063 Å in 4). However, despite the large steric effects, the planarity of C$_2$Al$_4$ is reasonably preserved (the distortion angle $\angle$ AlBRCCAlBR, $\delta = 169.7^\circ$), indicating the rigid C$_2$Al$_4$ cores. The t-Bu groups should bring large enough steric effects to protect the C$_2$Al$_4$ core. Similar strategy should be applicable to 5 and 6. We also found auxiliary supports for promising synthetic realization of such molecules. The C$_2$Cu$_4$ core in the synthesized tetranuclear acetylidocopper(ii)$^{36}$ has a structure similar to the C$_2$Al$_4$ cores.

In conclusion, we advanced our recently computed C$_2$Al$_4$ global minimum with dptCs to the new species which are promising for experimental access. In comparison with the isolated C$_2$Al$_4$, the new molecules have two advantages. Firstly, the eight non-bonding electrons in C$_2$Al$_4$ are more efficiently used for chemical bonding, which results in good thermodynamic stabilities. Secondly, the new molecules have “anchors” to attach protecting groups to hinder their possible high reactivity in the attempts to synthesize such molecules. We invite experimental realizations to move the phC chemistry forward.

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