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Stabilization of Planar Tetracoordinate Carbon^{1a}

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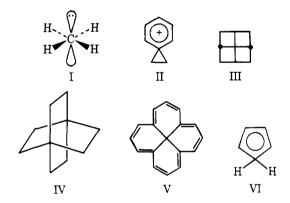
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Abstract: The energies of planar vs. tetrahedral geometries of tetracoordinate organic molecules have been surveyed by ab initio molecular orbital calculations. Because of their π acceptor and σ donor character, electropositive substituents, especially lithium, are particularly effective in stabilizing the planar arrangements selectively. Multiple substitution by such metals and by three-membered rings provides further stabilization, and 1,1-dilithiocyclopropane (XVII) and 3,3-dilithiocyclopropene (XVIII), inter alia, are actually calculated (RHF/STO-3G theory) to be more stable planar than tetrahedral.

van't Hoff and LeBel's proposal that tetracoordinate carbon prefers tetrahedral geometry² celebrated its centenary in 1974.3 No fundamental exception, e.g., an organic compound in which all four substituents lie in a plane, is known. It is not difficult to understand why this should be so. Singlet planar methane is sp² hybridized with a lone pair of electrons in the remaining p orbital (I).4 Consequently, only six electrons instead of eight are involved in bonding. Numerous theoretical calculations⁵ at different levels of sophistication have been applied to this problem and are summarized in Table I. The best available ab initio quantum mechanical calculations indicate D_{4h} singlet planar methane should be about 150 kcal/ mol less stable than the tetrahedral form.6 Since the bond dissociation energy of the C-H bond is only 104 kcal/mol⁷ D_{4h} planar methane would not be expected to exist under any circumstances. Figure 1 shows the molecular orbitals of planar methane.40

Nevertheless, chemists have long been fascinated with the possibility that compounds might be found in which a carbon either is planar or can undergo planar inversion with a sufficiently low barrier to permit experimental detection. The possibility that the stereomutation of cyclopropane might proceed via the twisting of one of the methylene groups through the carbon plane was considered^{8a} but has been rejected.^{8b} MINDO/2 calculations suggest that the phenonium ion (II) may have a low barrier to planar inversion, but this process has not been detected experimentally.⁹ trans-Fenestrane (III), ^{6a,10} [2.2.2.2]paddlane (IV), ¹¹ and aromatic molecules such as V^{4b} have been suggested as possibilities having planar carbons, but none of these have yet been synthesized.

III and IV represent the traditional "brute-force" approach to problems of this type; such systems will be highly strained if not prohibitively so. Instead of contriving against nature, it might be better to find ways preferentially to stabilize planar over tetrahedral geometries.



Previous investigations (Table II),^{4,12} particularly that of Hoffmann, Alder, and Wilcox, 4b have revealed how this might be done. For example, extended Hückel calculations (EHT) showed that the energy required to distort cyclopentadiene would be much reduced by the aromatic 6 π -electron character of the planar form (VI). σ-Donating electropositive substituents such as -SiH₃, -BH₂, and lithium should be effective in reducing the planar-tetrahedral energy difference.4b Since the "natural" angle in planar carbon derivatives is 90° instead of 109.5°, the introduction of small rings also should help to stabilize the planar forms since both angle strain and repulsive steric interactions between substituents would be reduced.9 This effect, along with π delocalization, contributes to the very low distortion energy (19^{12a} or 22^{12b} kcal/mol) calculated by MINDO/2 for the planar phenonium ion (II). Another approach often previously proposed4 is simply to remove two electrons thereby giving a six valence electron species. Unfortunately, CH₄²⁺, although calculated to prefer planarity strongly, is unstable toward dissociation into CH3+ and

Planar tetracoordinate carbon is a difficult problem to attack

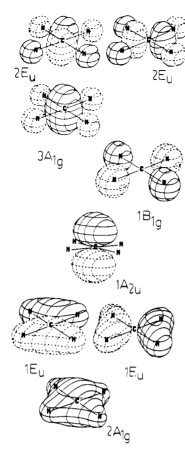


Figure 1. Valence molecular orbitals of planar methane (D_{4h}) arranged in order of increasing energy. The lower four are occupied in the singlet.

experimentally but it is easy computationally. Our approach has been to explore systematically by ab initio calculations the planar-tetrahedral energy difference in a variety of molecules. This has culminated in the discovery of relatively simple derivatives of methane that are indicated to prefer planar over tetrahedral geometries.

Computational Method

Several levels of sophistication were employed to determine the planar-tetrahedral energy differences. Most of the work was carried out using Hartree-Fock theory. For singlet states the spin-restricted form (RHF)^{15a} involving doubly occupied molecular orbitals was used but triplet states were handled by the spin-unrestricted version. 15b Initially the minimal STO-3G basis set¹⁴ was used in conjunction with standard geometries.¹⁶ Geometry optimization at the RHF/STO-3G level (or UHF/STO-3G for triplets) was then carried out, according to procedures described previously,17 on the smaller molecules and on those which were indicated to have small planar distortion energies. Calculated structures are given in Table III. Single calculations at the split valence RHF/4-31G or UHF/4-31G levels¹⁸ were then performed on the STO-3G minimized geometries. Total energies at these three levels are reported in Table IV. As can be seen from Tables V-VII, both geometry optimization and extension of the basis set uniformly reduce the energy differences. This arises since the "standard" models are appropriate for tetrahedral geometries but less so for planar structures largely because of significant π bonding or π antibonding between the substituent and the planar carbon atom; geometry optimization thus tends to lower the energy of the planar forms to a greater extent than the tetrahedral. Furthermore, since the planar carbon atoms have both σ and π bonds, they should benefit more from the increased flexibility

Table I. Calculated Energy Difference for Planar (D_{4h}) vs. Tetrahedral (T_d) Singlet Methane Using Several Molecular Orbital Methods (kcal/mol)

ΔE	Method or basis set	Ref
127	ЕНТ	4b
187	CNDO	4b
95	PNDO	5b
120	MINDO/3	а
250	Minimal STO	5a, d
240	RHF/STO-3G	5c, 6b
168	RHF/4-31G	5c, 6b
165	RHF/6-31G	6b
171	RHF/6-31G*	6b, <i>b</i>
166	RHF/6-31G**	6b, <i>b</i>
160	Double 5 with polarization	6a
157	RMP2/6-31G**	b
150	RHF/6-31G** with CI	6 b

a E. D. Jemmis, unpublished. b This work.

Table II. Semiempirical Calculations of Planar-Tetrahedral Energy Differences (kcal/mol)

Molecule	Method	ΔE
\triangleright	EHT	97a
	EHT	97 <i>b</i>
C(CN) ₄	EHT	79 <i>b</i>
⊕ H	EHT	67 <i>b</i>
$C(SiH_3)_4$ $C(BH_2)_4$	EHT EHT	67 <i>b</i> 42 <i>b</i>
(<u>+</u>)	ЕНТ	25 <i>b</i>
\bigcirc	MINDO/2	19¢
		22d

a Reference 9. b Reference 4b. c Reference 12a. d Reference 12b.

of the split valence basis set than do the tetrahedral geometries.

To confirm the reliability of these results, more rigorous studies were made in two cases. Methane has been treated most extensively. Reoptimization of the structure using the RHF/6-31G*19 level (which contains d-type polarization functions on carbon) yields a value for the barrier to planarity (171 kcal/mol) in close agreement with the RHF/4-31G result (168 kcal/mol) and a closely similar bond length.²⁰ Further extension of the basis set to include p functions on hydrogen (RHF/6-31G**) yields a value of 166 kcal/mol for the inversion barrier. Similarly a single calculation was performed at the RHF/6-31G* level for CH₂Li₂ at the STO-3G geometry.²⁰ The RHF/6-31G* barrier to inversion of 8 kcal/mol differs by only 2 kcal from the RHF/4-31G value, 10 kcal. A final check of the calculated structures was made by carrying out a partial geometry optimization of methyllithium (both C_{3v} and C_{2v}) using the extended RHF/4-31G level. This was done to ensure that the minimal STO-3G basis does not overestimate the role of the lithium 2p functions. Reoptimization of the C_{3v} structure resulted in an improvement in the RHF/4-31G total energy (-45.959 86 au) of only 0.2 kcal/ mol while the C-Li distance shortened 0.02 Å (2.008 to 1.990 Å) and the ∠HCLi angle changed 1.2° (112.6 to 111.4°). Reoptimization of the planar form gave a modestly larger lowering of 4.0 kcal/mol in the total energy (-46.898 30 au, RHF/4-31G) but a larger change in r(C-Li) (from 1.727 to 1.899 Å); ∠HCLi widens only slightly from 64.5 to 64.8°.

Similarly, RHF/4-31G reoptimization of tetrahedral CH₂Li₂ resulted in an improvement in the total energy (-53.77521 au) of only 0.4 kcal/mol and changes in the C-Li distance (1.924 to 1.966 Å) and LiCLi angle (119.8 to 115.0°) of 0.042 Å and 4.8°, respectively. Reoptimization of the planar form gave an energy decrease of 2.7 kcal/mol (-53.76231 au, RHF/4-31G), and an increase in the bond length (1.744 to 1.838 Å) of 0.094 Å and in the LiCLi angle (97.8 to 98.9°) of 1.1°. Based on these results, it seems that the 4-31G values listed in Tables V-VII are upper limits lying only a few kilocalories above the Hartree-Fock result.

The effect of electron correlation on the methane inversion barrier also was examined using restricted Møller-Plesset second-order perturbation theory²¹ applied to the 6-31G** wave function (this is designated RMP2/6-31G**). This method showed that inclusion of correlation energy led to a further lowering in the planar inversion barrier by 9 kcal/mol, from 166 to 157 kcal/mol (Table I).

An additional question is whether the electronic configuration depicted in I is truly the ground state. To test this the pair of electrons in the p orbital of planar methane was removed and placed in the electron deficient σ system (b_{1g} orbital). This results in an increase in the energy by 88 kcal (RHF/6-31G*) and abnormally long bond lengths of 1.237 Å. This result might not have been expected. Isoelectronic BH₄⁻ and valence isoelectronic SiH₄ prefer a vacant p orbital in their singlet planar ground states.²² The highest occupied b_{1g} orbital is then of d symmetry. The problem of whether the ground states are singlet or triplet is considered below.

Results and Discussion

Three effects should lower the planar distortion energy. Planar tetracoordinate carbon should be stabilized preferentially (1) by delocalizing the lone pair by π conjugation, ^{4b} (2) by providing more electron density to carbon by σ donation, ^{4b} and (3) by enforced reduction of the angle around the planar carbon atom by means of small rings.

Monosubstituted Methanes

Consider first the effect of a single substituent (Table V). Because of lone pair repulsion and high electronegativity, fluorine raises the planar-tetrahedral energy difference. On the other hand, the electropositive substituents, BH₂, BeH, and especially Li, lower the energy difference dramatically due to the simultaneous operation of σ -donating and π -accepting effects. Interestingly, the cyano group, a good π acceptor but also a good σ acceptor, is not particularly favorable.

Comparisons of all planar (e.g., VIIa) and perpendicular (e.g., VIIb) forms allow an assessment of the relative impor-

tance of σ vs. π effects with substituents such as BH₂, CH₂+, and NH₂. Delocalization of the carbon lone pair into the boron p orbital is precluded in perpendicular CH₃BH₂ (VIIb) resulting in an increase in the energy of the planar form by 26 kcal/mol (RHF/4-31G). This is a measure of the π effect. If BH₂ is replaced by isoelectronic CH₂+ the effect is even more dramatic, 115 kcal/mol (RHF/STO-3G). In the perpendicular forms (e.g., VIIb) only the σ effect should be operative (along with a modest amount of hyperconjugation), and the reduction of the planar-tetrahedral difference is 36 kcal/mol for a BH₂ substituent. Evidently the σ and π effects are of comparable significance in the influence of BH₂ on the planar-tetrahedral energy difference. Similarly, rotation of NH₂

in CH_3NH_2 (VIIIa,b) can eliminate the unfavorable C_{π} -N (lone pair) effect thereby lowering the planar distortion energy by 16 kcal/mol (RHF/4-31G).

$$\begin{array}{c} H \\ H \\ \downarrow \\ H \end{array} \longrightarrow \begin{array}{c} H \\ \downarrow \\ H \end{array} \longrightarrow \begin{array}{c} H \\ \downarrow \\ H \end{array} \longrightarrow \begin{array}{c} H \\ \downarrow \\ H \end{array}$$

$$\begin{array}{c} VIIIa \\ \end{array} \longrightarrow \begin{array}{c} VIIIb \\ \end{array}$$

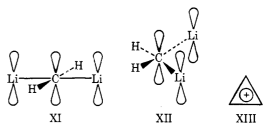
The calculated geometries (Table III) provide further insight into the electronic structure of the planar arrangements. Electropositive substituents which stabilize the planar form appear to be bridging a planar carbanion. Thus, in CH_3Li (IX) $\angle H_aCH_b$ is 115°, nearly the ideal 120° value. Destabilizing ligands such as fluorine (X) produce an opposite distortion

$$H_a$$
 H_b
 C
 H_a
 H_b
 C
 H_a
 H_a
 H_a
 C
 C
 C
 C

($\angle H_aCH_b$ is 63.3° in X); in these cases, H_b assumes the role of bridging ligand. Overlap populations indicate significant bonding between Li and H_a in IX and between H_a and H_b in X.

Polysubstituted Methanes

If one substituent lowers the distortion energy substantially, then two or more such substituents should be better. Although one does not normally think of disubstituted methanes as having cis and trans forms, this, of course, is possible if planar structures are adopted.²³ As illustrated in Table VI, if the substituents in question are metals, the cis planar isomers appear to be about 30 kcal/mol more stable than the trans. The electronic structure of trans-CH₂Li₂, with two π electrons in a linear three-center orbital (XI), is reminiscent of the allyl cation. The cis form, XII, is even better; the two π electrons



are delocalized in a cyclic arrangement, isoconjugate with the cyclopropenium ion (XIII). XII is "homoaromatic" in the sense that a π bond but not a σ bond exists between the two lithium atoms (cf. Figure 2). This is revealed by the Li…Li Mulliken overlap populations which are repulsive (-0.12) for the σ electrons, but bonding (+0.18) for the π . The 97.7° Li–C-Li bond angle in XII also indicates a balance between the attractive π and the repulsive σ Li…Li effects. The C-Li bond lengths in CH₂Li₂ decrease going from the tetrahedral² (1.924 Å) to the trans planar (1.807 Å) to the cis planar (1.744 Å) arrangements.

The third lithium atom in $CHLi_3$ produces a further, but modest, lowering of the tetrahedral-planar energy difference. The RHF/STO-3G minimized structure of the planar form reveals an even larger widening of the LiCLi angle to 101.7°. Because of geometric constraints, simultaneous angle enlargement of all LiCLi angles from 90° is impossible in planar CLi_4 . Evidently steric (repulsive σ) effects dominate whatever further favorable influences are produced by substitution of the fourth lithium atom, and the planar-tetrahedral energy

Molecule	Symmetry and structure	Optimized parameter	Molecule	Symmetry and structure	Optimized parameter
CH ₃ F (X)	C _{2v} planar	$r(C-H_a) = 1.124$	CH ₃ Na	C _{2v} planar	r(C-Na) = 2.131
3	24.	$r(C-H_b) = 1.136$	-	_,	$r(C-H_a) = 1.076$
		r(C-F) = 1.403 $\angle H_aCF = 116.7$			$r(C-H_b) = 1.053$ $\angle H_a CNa = 62.3$
CH ₃ OH (OH in plane)	C_{s}	r(C-O) = 1.473	CH ₃ CH ₂ ⁺	$C_{2\nu}$	r(C-C) = 1.318
ciigoii (Oii iii piiiio)	O.y	r(O-H) = 0.995	2 2		$r(C-H_a) = 1.114$
		$r(C-H_a \text{ cis}) = 1.118$			$r(C-H_b) = 1.184$
		$r(C-H_a \text{ trans}) = 1.107$ $r(C-H_b) = 1.136$			$r(C^+-H) = 1.094$ $\angle H_aCC = 113.4$
		$\angle COH = 101.1$			$\angle CC^{+} H = 121.3$
		$\angle OCH_a$ cis = 117.6	CH ₃ CH ₂ ⁺ (CH ₂ perpen-	$C_{2\nu}$	r(C-C) = 1.420
		$\angle OCH_a$ trans = 113.0	dicular)		$r(C-H_a) = 1.120$
CH ₃ OH (OH perpen-	C_{s}	$\angle OCH_b = 180.0 \text{ (ass.)}$ r(C-O) = 1.602			$r(C-H_b) = 1.144$ $r(C^+-H) = 1.128$
dicular)	-3	r(O-H) = 0.996			$\angle C^{\dagger}CH_{a} = 110.6$
		$r(C-H_a) = 1.117$			$\angle HC^{+}C = 124.0$
		$r(C-H_b) = 1.143$ $\angle COH = 84.9$	$CH_2(BeH)_2$	$C_{2\nu}$ tetrahedral	r(C-Be) = 1.666
		$\angle OCH_a = 86.6$			r(C-H) = 1.088 r(Be-H) = 1.290
		$\angle OCH_b = 180.0 \text{ (ass.)}$			∠HCH = 107.0
CH ₃ NH ₂ (VIIIa)	C_{S}	r(C-N) = 1.513			∠BeCBe = 113.9
		$r(C-H_a) = 1.104$ $r(C-H_b) = 1.131$	CU (DeU)	C planer	$\angle CBeH = 180.0$ r(C-Be) 1.557
		r(N-H) = 1.044	$CH_2(BeH)_2$	$C_{2\nu}$ planar	r(C-H) = 1.097
		$\angle NCH_a = 112.0$			r(Be-H) = 1.284
		ΔHNC = 104.7 ΔHNH = 101.7			\angle HCH = 105.3
CH ₃ NH ₂ (VIIIb)	C_{s}	r(C-N) = 1.438			∠BeCBe = 86.4 ∠BeBeH = 140.1
• • • •	a .	$r(C-H_a \text{ cis}) = 1.135$	$CH_2(BeH)_2$	D_{2h}	r(C-Be) = 1.580
		$r(C-H_a trans) =$	4	211	r(C-H) = 1.286
		1.113 r(N-H) = 1.036	HC(DoH)	C	r(Be-H) = 1.290
		$\angle NCH_a$ cis = 120.1	HC(BeH) ₃	C_{3v}	r(C-Be) = 1.644 r(C-H) = 1.093
		$\angle NCH_a$ trans = 116.0			\angle HCBe = 108.7
		∠HNC = 104.7 ∠HNH = 105.2	$HC(BeH)_3$	$C_{2\nu}$	$r(C-Be_a) = 1.616$
CH ₃ CH ₃	$C_{\mathcal{S}}$	r(C-C) = 1.507			$r(C-Be_b) = 1.546$ r(C-H) = 1.102
3	3	$r(C-H_a) = 1.106$			∠HCBe = 89.5
		$r(C-H_b) = 1.130$	C(BeH) ₄	D_{4h}	r(C-Be) = 1.622
		$r(C-H) = 1.091^{b}$ $\angle H_aCC = 113.9$	C(BeH) ₄	T_d	r(C-Be) = 1.630
		∠HCC = 112.2 ^b	cis-CH ₂ Li ₂ (XII)	$C_{2\nu}$ planar singlet	r(C-Li) = 1.744 r(C-H) = 1.110
CH ₃ BH ₂ (VIIa)	$C_{2 u}$ planar	$r(C-H_a) = 1.073$		Singlet	∠HCH = 101.4
		r(C-H) = 1.070 r(C-B) = 1.511			∠LiCLi = 97.7
		r(B-H) = 1.157	trans-CH ₂ Li ₂ (XI)	D_{2h} planar	r(C-Li) = 1.807 r(C-H) = 1.073
		$\angle H_aCB = 77.3$	CH ₂ Li ₂	singlet C_{2v} tetrahedral	r(C-H) = 1.073 r(C-H) = 1.091
CH ₃ BH ₂ (VIIb)	$C_{2\nu}$ planar	$\angle CBH = 120.9$ $r(C-H_a) = 1.156$	- 2 2	singlet	r(C-Li) = 1.923
Cli ₃ Dii ₂ (Viio)	C _{2v} planar	$r(C-H_b) = 1.051$			∠HCH = 106.7
		r(C-B) = 1.612	$\mathrm{CH_2Li_2}\;(\mathrm{XXIa})^b$	$C_{2\nu}$ tetrahedral	$\angle \text{LiCLi} = 119.8$ r(C-H) = 1.091
		r(B-H) = 1.166		triplet	r(C-Li) = 2.050
		∠H _a CB = 58.5 ∠CBH = 120.4			∠HCH = 105.2
CH ₃ BeH	C_{2v} planar	$r(C-H_a) = 1.113$. OTT 7 : (32327) h	0 1	∠LiCLi = 68.5
-		$r(C-H_b) = 1.064$	cis-CH ₂ Li ₂ (XXIb) ^b	$C_{2\nu}$ planar triplet	r(C-H) = 1.098 r(C-Li) = 1.999
		r(C-B) = 1.543 r(Be-H) = 1.280		tripict	∠HCH = 101.9
		$\angle HCBe = 66.29$			∠LiCLi = 68.9
CH ₃ Li (IX)	C_{2v} planar	$r(C-H_a) = 1.090$	CHLi,	C_{3v}	r(C-H) = 1.107
		$r(C-H_b) = 1.068$			r(C−Li) = 1.872 ∠HCLi = 104.8
		r(C–Li) = 1.727 ∠HCLi = 64.49	CHLi ₃	C_{2V} planar	r(C-H) = 1.135
CH ₃ CN	C_{3v}	r(C-H) = 1.088	•	2 , -	$r(C-Li_a) = 1.846$
-	2.	r(C-C) = 1.489			$r(C-Li_b) = 1.748$ \(\angle LiCH = 78.42\)
		$r(C \rightleftharpoons N) = 1.154$ $\angle HCC = 109.93$	CLi ₄	D_{4h}	r(C-Li) = 1.905
CH ₃ CN	$C_{2\nu}$ planar	$r(C-H_a) = 1.119$	CLi ₄	T_d	r(C-Li) = 1.859
,	~v •	$r(C-H_b) = 1.130$	CH ₂ Na ₂	C_{2v} tetrahedral	r(C-Na) = 2.050 r(C-H) = 1.09
		r(C-C) = 1.382 r(C-E) = 1.164			r(C-H) = 1.09 $\angle NaCNa = 123.0$
		$r(C \equiv N) = 1.164$ $\angle H_a CC = 114.68$			∠HCH = 109.47
CH ₃ Na	C_{3v}	r(C-Na) = 2.087	cis-CH ₂ Na ₂	C_{2V} planar	r(C-Na) = 1.946
	**	r(C-H) = 1.092			r(C-H) = 1.099 $\angle N_a C N_a = 92.9$
		∠HCNa = 114.3			'a 'a

Table III (Continued)

Molecule	Symmetry and structure	Optimized parameter	Molecule	Symmetry and structure	Optimized parameter
trans-CH ₂ Na ₂	D_{2h}	r(C-Na) = 2.012	Diboracyclopropane (XV)	$C_{2\nu}$ planar	r(C-B) = 1.514 r(C-H) = 1.085
cis-CLi ₂ F ₂	$C_{2\nu}$ planar	r(C-H) = 1.036 r(C-F) = 1.401			\angle HCH = 101.3
C13-CL1 ₂ 1 ₂	C _{2V} Planai	r(C-Li) = 1.695			$\angle BCB = 57.5$
		∠FCF = 99.50	Diboracyclopropane (XV)	$-C_{2\nu}$ tetrahedral	r(C-B) = 1.540
		4LiCLi = 117.1	Placetae, diopropune (111)	- 20	r(C-H) = 1.082
CLi ₂ F ₂	$C_{2\nu}$ tetrahedral	r(C-F) = 1.380			∠HCH = 110.53
02121 2	20 101141100141	r(C-Li) = 2.118			$\angle BCB = 63.1$
		$\angle FCF = 105.4$	2,3,4,5-Tetrabora-	$C_{2\nu}$ planar	r(C-B) = 1.527
		∠LiCLi = 114.6	spiropentane (XVI)	A7 A	$\angle BCB = 60.2$
Cyclopropene	$C_{2\nu}$ planar	$r(C_1 - C_3) = 1.693$	2,3,4,5-Tetrabora-	$C_{2\nu}$ tetrahedral	r(C-B) = 1.531
•		$r(C_3H) = 1.058$	spiropentane (XVI)	•	$\angle BCB = 61.2$
		$LC_1C_3C_2 = 43.0$	3,3-Dilithiocyclo-	$C_{2\nu}$ tetrahedral	$r(C_3 - Li) = 2.092$
		$\angle HC_3H = 124.4$	propene		$r(C_3 - C_1) = 1.498$
Cyclopropane	$C_{2\nu}$ planar	$r(C_1 - C_2) = 1.542$			\angle LiCLi = 102.6
		$r(C_1 - H) = 1.063$			$\angle C_1 C_3 C_2 = 50.8$
		$\angle C_2 C_1 C_3 = 54.4$	1,1-Dilithiocyclopropane	$C_{2\nu}$ planar	r(C-Li) = 1.733
	_	$\angle HC_1H = 110.8$	(XVIIc)		$r(C_1 - C_2) = 1.530$
Lithiocy clopro-	$C_{\mathcal{S}}$	$r(C_1-Li) = 1.702$			\angle LiCLi = 100.02
pane		$r(C_1 - C_2) = 1.496$			$\angle C_2 C_1 C_3 = 58.0$
		$r(C_1 - C_3) = r(C_1 - C_2)$	1,1-Dilithiocyclopropane	$C_{2\nu}$ tetrahedral	r(C-Li) = 1.952
		(ass.)	(XVIIa)		r(C-C) = 1.526
		$r(C_1 - H) = 1.068$			\angle LiCLi = 106.5
		$\angle C_2 C_1 C_3 = 60.9$			∠CCC = 58.6
		∠LiCH = 70.6 ∠CCLi = 93.3	3,3-Dilithio-1,2-diboracyclo-	$C_{2\nu}$ planar	r(C-Li) = 1.802
Tithia aralama	C total adval		propane (XIX)		r(C-B) = 1.485
Lithiocyclopro-	$C_{\mathcal{S}}$ tetrahedral	$r(C_1 - Li) = 1.961$			∠LiCLi = 129.5
pane		$r(C_1-C_2) = 1.513$ $r(C_1-H) = 1.084$	2250011 1200	C 1 1 :	∠BCB = 62.3
		$\angle C_2 C_1 C_3 = 59.0$	3,3-Dilithio-1,2-diboracyclo-	C_{2v} tetrahedral	r(C-Li) = 1.831
		$\angle \text{LiCH} = 117.8$	propane (XIX)		r(C-B) = 1.510
		$\angle \text{LiC}, C_2 = 123.9$			∠LiCLi = 123.7
		22.0102 123.3			$\angle BCB = 61.3$

a Bond lengths in $^{\circ}$ angles in degrees. See structures IX and X for specification of atoms. Unlisted parameters have standard values (ref 16). Exocyclic hydrogens lie in the plane perpendicular to and bisecting the ring angle. b Local $C_{3\nu}$ symmetry assumed for the tetrahedral methyl group which has dihedral angle H_{a} CCH of 90° . c UHF/STO-3G geometries for triplet states.

difference increases somewhat. This repulsive σ effect is less significant with BeH substituents, and a steady decrease in the planar-tetrahedral energy differences is noted in going from CH₃BeH (Table V) to C(BeH)₄ (Table VI).

The repulsive σ effect is revealed more dramatically in the case of $C(BH_2)_4$. The planar D_{4h} structure, XIVa, lies 54 kcal/mol higher in energy than the D_{2h} structure XIVb. This

is attributed to relief of the unfavorable interactions between hydrogen atoms. Alternatively, a "propeller" type distortion to the D_4 structure XIVc results in an energy intermediate between those of XIVa and XIVb. Evidently, a large amount of the favorable π delocalization must be sacrificed in order to overcome the unfavorable interligand steric effects.

Some second-row substituents, Na and Cl, were examined as well (Tables V, VI, and VIII). These elements demonstrated the same trends as were observed for the first-row substituents in that sodium lowered the inversion barrier in CH_3Na and CH_2Na_2 while Cl raised the barrier of CH_2LiCl relative to that of CH_3Li . However, sodium had a much smaller stabilizing effect than lithium presumably because of the diffuseness of its 3p orbitals. For the same reason, chlorine is not as destabilizing as fluorine since the unfavorable C_{π} -X (lone pair) (X

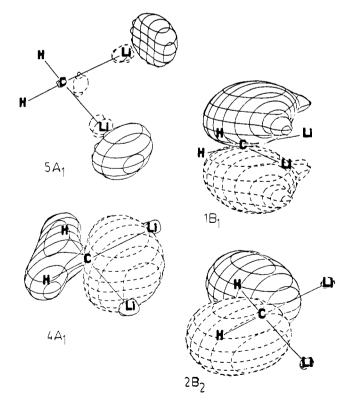


Figure 2. Four molecular orbitals of cis-planar $CH_2Li_2(C_{2v})$. The HOMO (1b₁) of the singlet shows the three-center, two-electron "homoaromatic" π bonding. The $5a_1$ MO is the LUMO in the singlet, but the HOMO in the triplet. Because of the diffuseness of the p orbitals on lithium, a contour of 0.06 au was employed.

Table IV. Total Energies (au) of Molecules in Tetrahedral² and Planar Arrangements

				STO-3G optim	nized geometry	
	RHF/STO-3G Star	ndard geometrya	RHF/S	TO-3G	RHF/	4-31G
Molecule	Tetrahedral	Planar	Tetrahedral	Planar	Tetrahedral	Planar
CH ₄ (I)	-39.726 86	-39.344 30	-39.726 86	-39.344 41	-40.13976	-39.871 47
CH ₃ F (X)	$-137.168 \ 36d$	-136.699 47	-137.169 06¢	-136.747 31	-138.856 86°	-138.514 99
CH ₃ OH (OH planar)	$-113.545 02^d$	-113.087 19	-113.545 98 <i>c</i>	-113.126 43	-114.865 25 c	-114.534 29
CH ₃ OH (OH perpendicular)	-94.030 43 <i>d</i>	-113.107 74 -93.581 69	-94.032 86 <i>c</i>	-113.148 61 - 92.619 96	-95.064 98¢	-114.546 03 - 94.748 87
CH ₃ NH ₂ (VIIIA) CH ₃ NH ₂ (VIIIB)	-34.030 43*	-93.600 24	-94.032 80	- 93.637 55	-93.004 965	- 94.774 74 - 94.774 74
CH ₃ CH ₃ (VIIID)	-78.306 18d	-77.909 53	-78.306 18 <i>c</i>	-77.921 26	-79.115 82¢	-78.837 69
CH ₃ BH ₂ (VIIA)	-64.666 50	-64.407 08	-64.667 69e	-64.421 52	-65.347 72	-65.179 65
CH ₃ BH ₂ (VIIB)		-64.328 26		-64.381 03		-65.138 08
CH ₃ BeH	-54.152 24	-53.929 26	$-54.153 \ 22f$	-53.993 63	-54.73349	-54.607 91
CH ₃ Li (IX)	-46.419 99	-46.247 88	$-46.421 59^f$	-46.338 05	-46.959 62	-46.891 92
CH ₃ CN	-130.27079	-129.902 23	-130.271 56	-129.925 61	-131.727 12	-131.475 02
CH ₃ Na	-198.937 38	-198.732 98	-198.940 68	-198.819 00	i do 104 06	i 55.056.24
CH ₃ CH ₂ ⁺ (CH ₂ planar)	-77.405 21	-77.204 26 -77.021 29	-77.408 06	-77.239 45	− 78.194 96	-77.076 34
CH ₃ CH ₂ ⁺ (CH ₂ perpendicular)		-77.021 29		-77.053 93		-77.925 46
cis-CH ₂ (BeH) ₂	-68.592 16	-68.476 71	-68.59299	-68.503 75	-69.338 31	-69.266 91
trans-CH ₂ (BeH) ₂	00.072 10	-68.431 08		-68.44252		-69.226 69
CH(BeH),	-83.050 72	-82.973 85	-83.053 18	-82.988 52		
C(BeH) ₄	-97.52643	b	-97.531 98	-97.480 43		
cis-CH ₂ Li ₂ (XII)	-53.131 15	-53.074 59	-53.134 07	-53.107 51	-53.774 54	-53.758 05
trans-CH ₂ Li ₂ (XI)		-53.030 64		-53.048 69		-53.698 81
cis-CH ₂ Li ₂ triplet (XXI)	t.	7	-53.152 50	-53.137 66	-53.799 94	-53.795 31
CHLi ₃	b b	$egin{array}{c} b \ b \end{array}$	-59.885 90 -66.666 68	-59.870 21 -66.631 93	-60.608 65 <i>0</i>	-60.596 82 <i>°</i>
CLi ₄ cis-CH ₂ Na ₂	-358.061 10	-357.958 14	-358.063 77	-357.978 44	-67.465 13 <i>°</i>	-67.438 90 <i>0</i> i
trans-CH ₂ Na ₂	330.001 10	-357.883 18	- 330.003 77	-357.891 73	i	i
cis-CHLi ₂ CN	-143.704.51	-143.651 63		307.071 70	•	·
trans-CHLi, CN		-143.60393				
cis-CLi ₂ F ₂ (XX)	-248.03279	-248.02057	-248.058 01	-248.07775	b	b
Cyclopropene	-114.393 25	-114.035 26	-114.401 168	-114.088 35	-115.641 68g	-115.40158
Cyclopropane	-115.660 37	-115.344 76	-155.666 16 <i>g</i>	-115.363 47	-116.883 50g	-116.659 23
Lithiocyclopropane	-122.356 56	-122.204 65	-122.364 59	-122.261 31	-123.698 98	-123.616 38
Diboracyclopropane (XV) 2,3,4,5-Tetrabora-	-88.342 75 $-137.020 04$	-88.260 18 -136.993 13	-88.344 33 -137.032 13	-88.289 85	-89.312 84	-89.281 81 -138.541 38
spiropentane (XVI)	-137.020 04	-130.993 13	-137.032 13	-137.015 75	-138.551 82	-136.341 36
1,1-Dilithiocyclopropane	-129.057 43	-129.03751	-129.059 86	-129.071 26	b	b
pane (XVII)	127.007	12,100, 01	227.007 00	1251011 20	Ü	v
3,3-Dilithiocyclopropene	-127.764 27	-127.777 52	-127.797 27	-127.812 11		
(XVIII)						
3,3-Dilithio-1,2-diboracy-	-101.876 82	-101.885 66	-101.888 69	-101.909 87	-103.061 93	-103.089 43
clopropane (XIX)	.00 .116 -01	100 115 001				
Cyclopentadiene (VI)	-190.446 53 <i>i</i>	-190.145 93j				
Phenonium ion (II)	$-304.231 42^{k}$	$-304.096 58^{k}$				
$C(BH_2)_4$ (XIVA)	-139.497 19	-139.277 62				
C(BH ₂) ₄ (XIVB)		-139.364 17 -139.323 89 n				
C(BH ₂) ₄ (XIVC)	-191.592 20	-139.325 89" -191.326 40				
Spiropentane CH ₂ LiF	-143.863 07	-143.644 17				
CH ₂ LiCl	-500.429 36	-500.222 77				
1-Lithio-1-fluoro-	-219.817 72	-219.60192				
cyclopropane						
C ₆ H ₇ ⁺	$-228.253 \ 16^{h}$	-228.034 99 ^h				
E H	-235.063 00 ^k	-234.988 86 ^k				
Li Li	-242.162 80 ^k	$-241.828 62^{k}$				
	-302.963 09 ^l	$-302.860 \ 70^{l}$				
Li Li	-197.044 75j	-197.168 61 ^m				
$ ightharpoons_{ ext{L}_{ ext{i}}}^{ ext{L}_{ ext{i}}}$	-203.877 03 <i>j</i>	$-203.902 88^{m}$				

^a See Table VII, footnote a. Parenthetical structural descriptions refer to orientations of substituents in the planar arrangements. ^b Convergence on the density matrix was not achieved. ^c W. A. Lathan, L. A. Curtiss, W. J. Hehre, J. B. Lisle, and J. A. Pople, *Prog. Phys. Org. Chem.*, 11, 175 (1974). ^d W. J. Hehre, R. Ditchfield, L. Radom, and J. A. Pople, J. Am. Chem. Soc., 92, 4796 (1970). ^e J. D. Dill, P. v. R. Schleyer, and J. A. Pople, *ibid.*, 97, 3402 (1975). ^f J. D. Dill, J. A. Pople, and P. v. R. Schleyer, to be published. ^g W. A. Lathan, L. Radom, P. C. Hariharan, W. J. Hehre, and J. A. Pople, *Top. Curr. Chem.*, 40, 1 (1973). ^h W. J. Hehre and J. A. Pople, J. Am. Chem. Soc., 94, 6901 (1972). ⁱ44-31G basis set not yet available for Na. ^j Experimental geometry for cyclopentadiene ring (L. H. Sharpen and V. W. Laurie, J. Chem. Phys., 43, 2765 (1965)). ^k Ring geometries from W. J. Hehre, J. Am. Chem. Soc., 94, 5919 (1972), for planar structure, cyclopropyl fragment rotated 90°. ^lC₆H₅ fragment from footnote k, cyclopropyl fragment from footnote g. ^m H. Preuss and G. Diercksen, Int. J. Quantum Chem., 1, 349 (1967). ⁿ Dihedral angle, φ(H,B,C,C) = 35°. ^o Convergence achieved using direct descent technique (R. Seeger and J. A. Pople, J. Chem. Phys., in press).

Table V. Calculated Planar – Tetrahedral² Energy Differences (kcal/mol) for Monosubstituted Methanes (RHF)

	Basis set		
Molecule	STO-3G (standard geometry)a	STO-3G 4-31C (STO-3G optimized geometry)	
CH ₄ (I)	240	240	168
CH ₂ F (X)	294	265	215
CH ₂ OH (OH planar)	287	263	208
CH ₃ OH (OH perpendicular)	274	249	200
CH ₂ NH ₂ (VIIIa)	281	259	198
CH ₂ NH ₂ (VIIIb)	269	248	182
CH ₂ CH ₃	250	243	175
CH ₃ BH ₂ (VIIa)	164	155	106
CH ₃ BH ₂ (VIIb)	213	180	132
CH ₃ BeH	140	100	79
CH ₃ Li (IX)	108	52	42 <i>b</i>
CH,CN	231	217	158
CH₃Na	124	76	c
CH ₃ CH ₃ + (CH ₂ coplanar)	128	106	74
CH ₃ CH ₂ + (CH ₂ perpendicular)	243	222	169

a Reference 16. b UHF/4-31G optimized structures have a difference of 38 kcal (see text). c 44-31G basis set not yet available for Na.

Table VI. Calculated Planar—Tetrahedral² Energy Differences (kcal/mol) for Polysubstituted Methanes (RHF)

	Basis set			
Molecule	STO-3G (standard geometry) ^a	STO-3G (STO-3G o		
cis-CH ₂ (BeH) ₂	72	56	45	
trans-CH, (BeH),	101	94	70	
CH(BeH),	48	41		
C(BeH)	c	32		
cis-CH ₂ Li ₂ (XII)	35	17	10^{b}	
trans-CH, Li, (XI)	63	54	47	
CHLi,	c	10	7	
CLi,	c	22	6	
cis-CH, Na,	65	59		
trans-CH, Na,	112	108		
cis-CHLi ₂ CN	33			
trans-CHLi2CN	63			
$CLi_2F_2(XX)$	8	-25	c	

^aReference 16. ^b8 kcal/mol both at 6-31G* (STO-3G geometries) and at 4-31G (4-31G geometries) (see text). ^cConvergence on the density matrix was not achieved.

= F, Cl) interaction is not as bad for the more diffuse chlorine 3p lone pair.

The high barriers to planarity of the carbenoids CH₂LiF and 1-lithio-1-fluorocyclopropane (Table VIII) support the recent findings of Hahnfield and Burton²⁴ on the stereochemical integrity of similar vinyl carbenoids. Contrary to an earlier report,²⁵ they found that these molecules did not isomerize in solution. Presumably, one possible mode of isomerization might have been a simple rotation of the LiCF group but this mechanism of rearrangement seems unlikely on the basis of our results.

Utilization of Small Rings

Three-membered rings overcome the steric problem. One can take advantage of such rings in two ways. The electropositive atoms can be incorporated into the ring structure, or they can be introduced as external substituents. Thus, XV meets the usual criteria for aromaticity in its planar arrangement, but the tetrahedral geometry is still preferred by 20 kcal/mol (RHF/4-31G). The same is true for XVI, a spiropentane analogue, but by only 6 kcal/mol (RHF/4-31G).

Table VII. Calculated Planar—Tetrahedral² Energy Differences (kcal/mol) for Three-Membered Ring Compounds (RHF)

	Basis set	
STO-3G (standard geometry) ^a	STO-3G (STO-3G o geome	
225	196	151
198	190	141
95	65	52
52	34	20
17	10	6
13	-7	b
-8	$(-10)^d$	b
-6	-13	-17
	(standard geometry)a 225 198 95 52 17 13 -8	STO-3G (standard geometry) ^a STO-3G of (STO-3G of Geometry) ^a geometry) ^a geometry) ^a 196 198 190 95 65 52 34 17 10 13 -7 -8 (-10) ^a

a Reference 16; for planar molecules the standard bond angle is 90° except when a ring is present. In these cases the remaining substituents trisect the external angle. b Convergence on the density matrix was not achieved. Cone carbon planar. Incomplete optimization.

Table VIII. Calculated Planar—Tetrahedral² Energy Differences (RHF/STO-3G, kcal/mol) for Some Larger Systems at Standard Geometries

189
107
85
84
138
113
167
137
130
135
137
61
21
102
77
16

^a See Table VII, footnote a, and Table III for the geometries of individual molecules. ^b Central carbon planar.

1,1-Dilithiocyclopropane (XVII) achieves our objective; the planar form is calculated even at the RHF/STO-3G level to be the energy minimum! 3,3-Dilithiocyclopropene (XVIII), the rather exotic compound XIX, and the bis-carbenoid CLi_2F_2 (XX) are even more favorable in their preference for planar geometries.

Triplet States

An obvious additional question, emphasized by Murrell, 5e is the possibility of a triplet ground state. UHF/6-31G** calculations by Shavitt et al.6b indicate that triplet planar methane has an energy 2.5 kcal/mol lower than singlet planar methane. However, the problem of assessing triplet-singlet energy differences by single-determinant calculations is well known.²⁶ The stability of states of higher spin multiplicity is overestimated. For example, the stability of the triplet state of CH₂ is overestimated by $14 \pm 6 \text{ kcal/mol}^{27}$ at the Hartree-Fock level. Therefore, it is not surprising that when configuration interaction is employed singlet methane becomes the more stable planar configuration by 4 kcal/mol.6b We have calculated the singlet-triplet difference for CH₂Li₂. The total energy of the planar triplet (UHF/4-31G) lies 23.4 kcal/mol below that of the planar singlet (RHF/4-31G). The tetrahedral triplet (UHF/4-31G) is only 15.9 kcal/mol below the tetrahedral singlet (RHF/4-31G). The planar distortion energy of the triplet is then only 2.9 kcal/mol (UHF/4-31G). In view of the overestimation of the stability of triplet methylene, it is difficult to assign the ground state configuration of CH₂Li₂ but the two states should be close in energy.

The structures of the two triplets, XXIa,b, show marked differences from the corresponding singlet geometries. For both

triplets an electron is removed from a LiCLi orbital of π type (symmetry b_1) and placed in a Li···Li σ bonding orbital (symmetry a_1). This results in a marked reduction of the \angle LiCLi angles (to ca. 69° in both XXIa and XXIb); three-membered rings are formed. The C-Li bond lengthenings (Table III) reflect the removal of an electron from the π bonding orbital. The presence of low-lying triplet configurations may contribute to the tendency of polylithium compounds to associate, 28 but nothing is known about this at present.

Geometries of Lower Symmetry

There remains the problem of determining the most stable geometry. Orientations other than T_d and D_{4h} are certainly possible for methane as are corresponding structures for substituted methanes. Indeed, Shavitt's work6b indicates that a square-pyramidal (C_{4v}) geometry is preferred over square planar (D_{4h}) for methane suggesting that molecules such as cis, cis, cis, cis-fenestrane 10b would undergo a similar distortion as do several known carboranes.²⁹ However, complete relaxation of all symmetry constraints would result in a prohibitive computational burden and was not generally pursued here. In two cases we did carry out a somewhat more complete search (at the RHF/STO-3G level) to probe the possibility that a structure intermediate between the "tetrahedral" C_{2v} and planar C_{2v} structures might be even lower in energy. For the case of CF_2Li_2 , a form of C_2 symmetry, obtained by rotating the FCF plane in the cis planar form by 20° about the angle bisector, was found to lie 0.7 kcal/mol lower in energy than the planar structure. In contrast, a 5° rotation of the CLi2 group in planar 1,1-dilithiocyclopropane (XVIIc) resulted in a higher energy.

Two promising systems, 5-lithiocyclopentadiene and 5,5-dilithiocyclopentadiene (XXIIa and XXIIb), received no further attention since calculations at fixed geometries³⁰ showed that these molecules prefer the unrelated bridged structure XXIIa,b.

Proposed Experimental Systems for Study

After a century of tetrahedral carbon, these results are startling and invite experimental verification. For molecules in which the tetrahedral structures are more stable, but the tetrahedral-planar energy difference is low, stereomutation should be observable.³¹ Appropriately substituted phenonium ions are a possibility, but our ab initio results on II (Table VIII), although only with standard geometries, do not confirm the low MINDO/2 distortion energies^{12,32} and preliminary experimental work failed to detect planar inversion.⁹ A variety of alkyllithium compounds, including neohexyllithium³³ and the cyclopropyllithium derivative (XXIII),³⁴ are known to

stereomutate in solution.²⁶ A dissociation-recombination mechanism involving ions or ion pairs has generally been assumed.³³ Electron donor solvents may well favor this mechanism, but we suggest that planar inversion should receive serious consideration as an alternative in nonpolar solvents especially since the distortion energy for methyllithium is calculated to be only 38 kcal/mol (RHF/4-31G optimized structures). Furthermore the structures we have calculated for planar alkyllithium compounds resemble trigonal carbanions bridged by lithium cations with significant bonding, not merely electrostatic attraction, between lithium and carbon as well as between lithium and the other adjacent atoms, especially hydrogen.³⁵ Stereomutation of alkyllithium derivatives may also occur in associated species, but we have not yet considered such possibilities computationally.

Evidence for the existence of many di- and polylithium compounds, including CH₂Li₂,³⁶ CLi₄,³⁷ and the 1,1-di- lithiocyclopropane derivative (XXIV),³⁸ has been reported,^{26,39} but their structures are not known experimentally. Our results indicate that molecules of this type, at least as isolated species in the gas phase, are likely to be planar or to have low planar distortion energies. Although the experimental verification of these predictions may be complicated by the tendency of lithium compounds to be associated or to be solvated in electron-donor solvents,²⁶ efforts to overcome these problems should be rewarding.

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- Exact tetrahedral geometry (T_d symmetry) is seldom encountered, as it is possible only when all four substituents are the same. As deviations from 109.5° bond angles are generally small, "tetrahedral" is often employed as we do here in a loose sense to describe the preferred bonding arrangement in saturated organic molecules.
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