Low-Lying Electronic States of Unsaturated Carbenes. Comparison with Methylene

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Abstract: The lowest singlet and triplet electronic states of methylene, H2C:, vinylidene, H2CC:, and vinylidenecarbene, H2CCC:, are studied theoretically at the spin-unrestricted Hartree-Fock (HF) and spin-unrestricted many-body perturbation theory (MBPT) levels of approximation. Calculations at both levels predict a triplet ground state for methylene and singlet ground states for vinylidene and vinylidenecarbene. MBPT singlet-triplet energy splittings are found to be ~13.1 kcal/mol for methylene, 51.1 kcal/mol for vinylidene, and 48.7 kcal/mol for vinylidenecarbene. A comparison of the effects of correlation on electronic structure is made between the parent carbene, methylene, and each of these unsaturated carbenes. From electronic structure trends observed in H2C:, H2CC:, and H2CCC:, singlet ground states are predicted for the extended unsaturated carbenes H2CCC:, and H2CCC, the next members of the series. The importance of d-type carbon polarization functions in basis sets used for the calculation of electronic structure properties in unsaturated carbenes is demonstrated by comparison with results in which d functions are not included in the basis.

Introduction

In recent years, increasing attention has been focused on reactions in which unsaturated carbenes appear as intermediates.2,3 However, little is known about the "isolated-molecule" properties of these highly reactive species, since direct spectroscopic observation has not yet proven to be generally feasible. Because the two lowest carbene electronic states that one is interested in probing are of different spin symmetry, the use of conventional optical spectroscopy is not practical. To the best of our knowledge, the photodetachment of H2C:− by Zittel et al.1 to yield 1A1 H2C: and 3B1 H2C: is the only quantitative gas-phase experimental investigation of the singlet-triplet splitting in a carbene.5 Several earlier studies have been reported, however, in which the methylene singlet-triplet splitting was estimated from the photolysis of ketene, CH3CO.6,7 Analogous experiments have not yet been performed for any unsaturated carbene.

Although much valuable information about the electronic states of other carbenes has been inferred from the nature of the chemical reactions they undergo2-3 (e.g., one-step stereospecific addition to olefins for the singlets, two-step nonspecific reactions for the triplets), the validity of such inferences often depends upon assumptions concerning the rate at which the nascent carbene is relaxed to its ground state. Moreover, the results of such condensed-phase experiments contain the effects of the solvent on the carbene's electronic states. In light of the above experimental difficulties, we believe that it is of great importance to acquire a more fundamental understanding of those aspects of the carbene's geometrical and electronic structure governing the relative stability of the lowest singlet and triplet states in the "gas-phase" species. Because of the considerable difficulty that arises in experimental studies of these highly reactive intermediates, state of the art quantum chemical calculations provide the only practical and reliable access to most of the spectroscopic, bonding, and geometrical information bearing upon the chemical behavior of these carbenes. The potential contributions of quantum theoretical calculations to carbene chemistry have already become rather clear. Since Mulliken's pioneering theoretical work in 1932,8 numerous authors have investigated the important methylene problem by means of semiempirical and, more recently, ab initio9-28 theoretical methods. In particular, Staemmler,15,16 Harrison,10,20 Bender et al.,17,23,24 Schaefer et al.,17-26 Bauschlicher et al.,17,23,24,26,29 and Harding and Goddard27 have carried out careful ab initio studies on methylene within the last 5 years that have had substantial impact on the area. These studies have served to clarify the interpretation of experimental data (e.g., the geometry of the triplet ground state) and to place in proper perspective the body of earlier ab initio and semiempirical work in which the highly correlated 1A1 state was not well described owing to deficiencies in the choice of basis set and/or configuration space. However, only a few semiempirical20,21 and ab initio22-24 calculations have been performed on vinylidene, H2C=C:, the simplest unsaturated carbene, and to our knowledge, only one HF-level ab initio theoretical study has been done on vinylidenecarbene, H2C=C=C:, the prototypical system for the class of RR'C=C= intermediates whose reactions constitute an active and growing area of experimental focus in carbene chemistry.2,3

In this paper, we explore the electronic structure of vinylidene (2) and vinylidenecarbene (3) at several levels of theoretical approximation, making comparisons, where appropriate, with methylene (1). Our study concentrates on two properties of particular interest in carbene chemistry: the spin multiplicity of the electronic ground state and the energy splitting between this state and the first excited state. These properties are clearly very relevant to an understanding of the chemical behavior of unsaturated carbenes. For example, reaction channel choice must accommodate the spin multiplicity of the carbene intermediate without violating quantum mechanical state symmetry constraints. Similarly, the susceptibility of such reactions to thermal or photochemical alteration is determined in part by the energy gap between the ground and first excited states of the intermediate carbene species.

Computational Details

A. Geometry. Calculations on 1 were carried out at the geometries of the 1A1 and 3B1 potential surface minima as determined by Bauschlicher and Shavitt (1A1, θHCH = 102.4°, RCH = 1.116 Å; 3B1, θHCH = 132.4°, RCH = 1.082 Å).29

Spin-unrestricted HF and correlated calculations on 2 were performed at the geometry of the 3B2 potential surface minimum reported by Davis, Goddard, and Harding (C2v symmetry, RCH = 1.076 Å, θHCH = 116.6°, RCC = 1.36 Å).35 It is reasoned that the HCH angle in 2 should be similar to the
ethylene HCH angle. HF-level calculations on this system were also carried out with $R_{CC} = 1.35 \text{ Å}$, the geometry these authors identify as the $^1\text{A}_1$ potential surface minimum. Our HF-level calculations indicate that changes due to the 0.01 Å C–C bond distance change are small. Hence, we have restricted our correlated calculations to the Davis, Goddard, and Harding triplet minimum energy geometry. For carbene 3, all calculations were carried out with the same C–H distances and HCH angle used for 2. We make the reasonable assumption that these parameters would be influenced only slightly by the attachment of another carbon atom at the opposite end of carbene 2. The C–C bond distances were shortened to 1.32 Å in 3 to correspond more closely with the C–C distance of 1.31 Å reported by Herzberg for the structurally similar species allene, $\text{H}_2\text{C} = \text{C} = \text{C} = \text{H}_2$.36

It should be stressed that the singlet–triplet energy splittings we report for carbens 2 and 3 are vertical rather than adiabatic splittings. Choice of geometries at which to perform ab initio calculations on larger many-electron systems such as 2 or 3 necessarily involves a compromise between computational expense and detailed knowledge of the potential energy surfaces of the states of interest. While our geometries were selected with careful regard for relevant experimental and theoretical information, the picture of the singlet and triplet potential energy surfaces we present is extremely limited. The arguments advanced by Moule and Walsh with regard to formaldehyde lend support to our assumption of planar $C_2\text{v}$ symmetry in the $^1\text{A}_1$ and $^3\text{B}_2$ states of 2 and 3.37 A more detailed study of these states, which will consider variations of C–C bond coordinates and the HCH angular coordinate, is now in progress and will be reported in a subsequent paper.

B. Basis Sets. Three types of basis sets were used in the ab initio calculations. The Dunning (4s2p/2s) = (C/H) contraction38 of Huzinaga’s (9s4p/4s) primitive Gaussian basis39 served as the reference basis set in this work. Other basis sets were derived from this by addition of polarization functions, known to be important in theoretical descriptions of $\text{H}_2\text{O}$, $\text{CH}_2\text{N}$, and $\text{CCl}_2$ to the carbene or terminal carbon atoms in 2 and 3. The need for better description of the terminal carbon atom is suggested by the localization of the nonbonding valence electrons in this region of the unsaturated carbene. In carbene 2, a basis set containing d functions on both carbon atoms, a duplication of the Davis, Goddard, and Harding basis set,33 was also used. Comparison between HF-level results on 2 employing one and two d function basis sets suggests that restriction of the d function to the terminal carbon is a reasonable compromise between basis set size and basis flexibility (e.g., $^1\text{A}_1 \rightarrow ^3\text{B}_2$ HF-level splitting in 2 is 11.1 kcal/mol with the (4s2p,4s2p/2s) basis, 17.6 kcal/mol with the (4s2p,4s2p,4s2p/2s) basis, and 15.7 kcal/mol with the (4s2p,4s2p,4s2p/2s) basis. $^1\text{A}_1$ HF-level energies for the three basis sets are –76.7401, –76.7592, and –76.7634 hartrees, respectively). It is seen that adding a d function on the carbon carbene stabilizes the $^1\text{A}_1$ HF state by 6.5 kcal/mol relative to the $^3\text{B}_2$ HF state. By including a methane carbon d function as well, the $^3\text{B}_2$ HF state is stabilized by another 1.9 kcal/mol, which reduces the $^1\text{A}_1$ stabilization for this basis to 4.6 kcal/mol. Although a carbene carbon d function alone slightly overestimates the $^1\text{A}_1$ HF state, its presence in the basis yields a much better overall picture of the singlet–triplet splitting. For our correlated studies of 1, 2, and 3, we employ the (4s2p/2s), (4s2p,4s2p/2s), and (4s2p,4s2p,4s2p/2s) basis sets, respectively. A d function exponent of 0.6769 was used in all of our calculations.33

C. Ab Initio Methods. Symmetry-adapted atomic integrals for use in the HF and correlated calculations were generated by the MOLECULE integral program.40 This is a component of the PROPAGATOR program system for the calculation of molecular electronic structure, which uses many-body tech-

### Table I. Methylene (H₂C) Calculations<sup>d,e</sup>

<table>
<thead>
<tr>
<th>Method</th>
<th>$^1\text{A}_1$</th>
<th>$^3\text{B}_1$</th>
<th>$\Delta (^1\text{A}_1 \rightarrow ^3\text{B}_1)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HF</td>
<td>–38.8766</td>
<td>–38.9221</td>
<td>0.0455</td>
</tr>
<tr>
<td>MBPT-4</td>
<td>–38.9967</td>
<td>–39.0201</td>
<td>0.0234</td>
</tr>
<tr>
<td>MBPT-[3-2]</td>
<td>–39.0020</td>
<td>–39.0221</td>
<td>0.0201</td>
</tr>
<tr>
<td>MBPT-[3-3]</td>
<td>–39.0012</td>
<td>–39.0220</td>
<td>0.0208</td>
</tr>
<tr>
<td>MBPT-8</td>
<td>–39.0023</td>
<td>–39.0223</td>
<td>0.0198</td>
</tr>
</tbody>
</table>

<sup>d</sup> hartrees (1 hartree = 627.5 kcal/mol).<sup>e</sup> kcal/mol. * (2S + 1) = 3.01 UHF. d (4s2p/2s) basis set, mixed hydrogen exponential scale factors, $s_1 = (1.2,1.0,1.0), s_2 = (1.0)$.<sup>f</sup> Geometries: $^1\text{A}_1$, $R_{\text{HCH}} = 102.4^\circ$, $R_{\text{CH}} = 1.116$ Å; $^3\text{B}_2$, $R_{\text{HCH}} = 132.4^\circ$, $R_{\text{CH}} = 1.082$ Å.

### Table II. Vinylidene (H₂CC) Calculations<sup>d,e</sup>

<table>
<thead>
<tr>
<th>Method</th>
<th>$^1\text{A}_1$</th>
<th>$^3\text{B}_1$</th>
<th>$\Delta (^1\text{A}_1 \rightarrow ^3\text{B}_1)$</th>
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<tr>
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<td>–76.7634</td>
<td>–76.7394</td>
<td>0.0250</td>
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<td>MBPT-4</td>
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<td>–76.9354</td>
<td>0.0792</td>
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<td>MBPT-[1-1]</td>
<td>–77.0193</td>
<td>–76.9375</td>
<td>0.0818</td>
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<tr>
<td>MBPT-8</td>
<td>–77.0207</td>
<td>–76.9393</td>
<td>0.0814</td>
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<tr>
<td>MBPT-[3-3]</td>
<td>–77.0215</td>
<td>–76.9401</td>
<td>0.0814</td>
</tr>
<tr>
<td>DECI-RSPT-4</td>
<td>–77.0016</td>
<td>–76.9295</td>
<td>0.0721</td>
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</table>

<sup>d</sup> hartrees (1 hartree = 627.5 kcal/mol).<sup>e</sup> kcal/mol. * (2S + 1) = 3.27 UHF. d (4s2p,4s2p/2s) basis set, hydrogen exponential scale factor = 1.0. Geometries: $^1\text{A}_1$, $R_{\text{HCH}} = 116.6^\circ$, $R_{\text{CH}} = 1.076$ Å; $R_{\text{CC}} = 1.36$ Å.

### Table III. Vinylidenecarbene (H₂CCH) Calculations<sup>d,e</sup>

<table>
<thead>
<tr>
<th>Method</th>
<th>$^1\text{A}_1$</th>
<th>$^3\text{B}_1$</th>
<th>$\Delta (^1\text{A}_1 \rightarrow ^3\text{B}_1)$</th>
</tr>
</thead>
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<tr>
<td>HF</td>
<td>–114.5800</td>
<td>–114.5761</td>
<td>0.0039</td>
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<td>MBPT-4</td>
<td>–114.8799</td>
<td>–114.8046</td>
<td>0.0753</td>
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<td>MBPT-[1-1]</td>
<td>–114.8929</td>
<td>–114.8081</td>
<td>0.0848</td>
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<tr>
<td>MBPT-8</td>
<td>–114.8885</td>
<td>–114.8109</td>
<td>0.0776</td>
</tr>
<tr>
<td>MBPT-[3-3]</td>
<td>–114.8897</td>
<td>–114.8138</td>
<td>0.0759</td>
</tr>
<tr>
<td>DECI-RSPT-4</td>
<td>–114.8554</td>
<td>–114.7975</td>
<td>0.0579</td>
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<sup>d</sup> hartrees (1 hartree = 627.5 kcal/mol).<sup>e</sup> kcal/mol. * (2S + 1) = 1.40 UHF. d (4s2p,4s2p,4s2p/2s) basis set, hydrogen exponential scale factor = 1.0. Geometries: $^1\text{A}_1$, $R_{\text{HCH}} = 116.6^\circ$, $R_{\text{CH}} = 1.076$ Å; $R_{\text{CC}} = 1.32$ Å.

### References

41-44 Developed and implemented by two of us (G.P. and R.J.B.). Singlet HF calculations were restricted to be eigenfunctions of spin and spatial symmetry. HF calculations on the triplet states were performed using the spin-unrestricted formalism (UHF) in which spatial parts of orbitals of different spin are allowed to be different. However, each spin orbital was required to be of pure spatial symmetry. For example, orbitals of symmetry $\text{a}_1$ were not allowed to mix with orbitals of other symmetries such as $\text{b}_2$. Since it is well known that UHF-level calculations are not exact eigenfunctions of spin, expectation values of the spin multiplicity (2S + 1) were calculated for the triplet states at the UHF level and are reported in Tables I–III.

Correlated calculations were obtained by adding to the HF total energies the many body perturbation theory (MBPT) correlation corrections that arise from double excitation type diagrams (DEMBPT) through eighth order.41,42 This double excitation method is particularly appropriate for treating correlation in low-lying $^1\text{A}_1$ carbene states where $\pi^2 \rightarrow \pi^2$ double-excitation configurations are known to be important. For the triplet states, a spin-unrestricted MBPT formalism was used.45 Since higher order corrections to the average spin multiplicity are introduced by the correlation, this method provides a better approximation to the triplet multiplicity than the UHF. The DEMBPT model is similar in conceptual simplicity to the established double-excitation configuration interaction method (DECI), but avoids the systematic "size-
Table IV. HF and Correlated 1A1-3B1 Energy Splittings, Correlation Energies, and Correlation Energy Differences (kcal/mol)\(^a\)

<table>
<thead>
<tr>
<th>carbene</th>
<th>(\Delta E_{HF})</th>
<th>(\epsilon_1(A_1))</th>
<th>(\epsilon_2(3B_1))</th>
<th>(\Delta)</th>
<th>(\Delta E_{HF} + \Delta\epsilon_1 = \Delta E_c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHC:</td>
<td>(-28.6), -25.0(^a)</td>
<td>(78.2), 95.3(^a), 13.9(^b), -24.8(^c)</td>
<td>(62.7), 80.9(^c), 0.0(^b), (15.5), 14.4(^d), 13.9(^b), (-13.1), -10.6(^c), -11.1(^b), -10.8(^c)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HCC:</td>
<td>(15.7), 31.0, 27.2(^d)</td>
<td>(161.5), 70.7, 103.2(^d)</td>
<td>(126.1), 55.8, 98.0(^e), (35.4), 14.9, 5.2(^f)</td>
<td>(51.1), 45.9, 32.4(^f)</td>
<td></td>
</tr>
<tr>
<td>HCCC:</td>
<td>(2.4), -1.5(^f)</td>
<td>(193.6), (147.3)</td>
<td>(46.3)</td>
<td>(48.7)</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Reference 29, (6s4p2d/3s2p) basis set. UHF triplet, multiconfiguration correlation. \(^b\) Reference 29, (6s4p2d/3s2p) basis set. UHF triplet, two-configuration singlet. \(^c\) Reference 27, 5s3p2d/2s basis set. UHF triplet, multiconfiguration correlation. \(^d\) Reference 33, (4s2p4d2s/2s) basis set. RHF triplet, multiconfiguration correlation. \(^e\) Reference 34, (4s2p4d2s/2s) basis set. RHF triplet, limited correlation. \(^f\) Reference 35, (6-31G*) polarized basis set. UHF triplet. Our HF and MBPT-8 results appear in parentheses. Basis sets used in our work are 1, (4s2p4d2s/2s); 2, (4s2p4d2s/2s); 3, (4s2p4s2p4s2p/2s). Our HF-level splittings are taken with respect to a UHF triplet.

A. Methylene. Our discussion of the low-lying states of unsaturated carbene 2 and 3 is conveniently prefaced by a simple orbital description of the two lowest states of carbene 1. Following the IUPAC suggestion for molecules with \(C_2\) symmetry,\(^37\) the \(z\) axis is chosen as the molecular symmetry axis and the hydrogens are assumed to lie in the \(yz\) plane. With this convention, \(\sigma\) orbitals along the \(z\) axis have \(a_1\) symmetry, \(\pi\) orbitals perpendicular to the molecular plane have \(b_1\) symmetry, \(\pi\) orbitals parallel to the molecular plane and perpendicular to the \(z\) axis have \(b_2\) symmetry, and \(C-H\) bonding orbitals have either \(a_1\) or \(b_2\) symmetry. We retain this convention for carbene 2 and 3.

The simple molecular orbital (MO) picture of the \(3B_1\) ground state of carbene 1 is shown in A. It consists of a filled carbon 1s atomic orbital with \(a_1\) symmetry, two \(C-H\) bonding orbitals of \(a_1\) and \(b_2\) symmetry, and a nonbonding valence orbital occupancy designated by \(\sigma\), or \(a_1\).

\[1B_1)_{1HxC} \cong [1a_1^22a_2^21b_2^2] 3a_11b_1 = [\text{core}] a_1b_1 \]

The first excited state of the \(1A_1\) state, can be described as a mixture of the two configurations B and C with the \(B\) configuration and the \(C\) secondary but important \(3a_1^2 \rightarrow 1b_2^2\) double-excitation configuration of B.

\[|1A_1\rangle_{1HxC} \approx C_B[\text{core}] a_1^2 + C_C[\text{core}] b_1^2 \]

In eq 2, we have \([C_B]|C_C| \approx 1\) (e.g., \(C_B = 0.9617\) and \(C_C = 0.943\) in the CI study of O’Neil, Schaefer, and Bender). Several authors have pointed out that the two-configuration description of the \(1A_1\) state is necessary to ensure treatment of the \(a_1\) and \(b_2\) valence orbitals on an equivalent footing at a \"HF\" level of approximation.\(^{10,12,13,25,26,28}\) In the \(1B_1\) ground state, the single configuration A suffices, since the valence orbital occupancy includes both of these orbitals.

It should be noted that the two-configuration valence orbital description of the \(1A_1\) state and the single-configuration function for the \(1B_1\) state, Meadows and Schaefer and Bauschlicher and Shavitt\(^25\) obtained adiabatic \(1A_1-3B_1\) energy splittings of \(-10.9\) kcal/mol. However, these values are in almost exact agreement with the best multiconfiguration results of Bauschlicher and Shavitt,\(^29\) and independently, Harding and Goddard,\(^27\) at \(-10.6\) kcal/mol. From this, we conclude that the \(1A_1\) and \(3B_1\) core-electron correlation energies are nearly identical, which allows the simple orbital descriptions (eq 1 and 2) to model accurately the distinctive valence characteristics of these states.

Single-configuration HF calculations on both states of 1 (i.e., \(C_B = 1, C_C = 0\) in eq 2) by Meadows and Schaefer\(^25\) and Bauschlicher and Shavitt\(^29\) gave splittings of \(-24.8\) and \(-25.0\) kcal/mol, respectively. Thus, a correlation contribution of \([24.8 - 10.9]^2 = [25.0 - 11.0]^2 = 13.9\) kcal/mol can be ascribed to the double-excitation configuration C of the \(1A_1\) state. While multiconfiguration correlation significantly lowers the energy of each state (e.g., with respect to single-configuration HF values, Bauschlicher and Shavitt report that CI lowers the \(1A_1\) energy by 95.3 kcal/mol and the \(1B_1\) energy by 80.9 kcal/mol, almost all of the [95.3 – 80.9] = [25.0 - 10.6] = 14.4 kcal/mol HF-CI differential decrease can be attributed to the presence of configuration C in the multiconfiguration \(1A_1\) state, since this value is within 0.5 kcal/mol of the 13.9 kcal/mol one- vs. two-configuration differential decrease calculated with the same basis set and geometry.\(^29\) Thus, including the double-excitation configuration C in the \(1A_1\) state description, either at the two- or multiconfiguration level, lowers the \(1A_1\) excited state energy about 14 kcal/mol relative to the corresponding \(3B_1\) ground-state energy, hence diminishing the \(1A_1-3B_1\) splitting by this amount as well. These results are summarized in Table IV.

To test the applicability of the MBPT method as a model for carbene electronic structure within our choice of basis set, we performed calculations on the \(1A_1\) and \(3B_1\) states of 1 at the
minimum energy geometries reported by Bauschlicher and Shavitt. These results are presented in Table I. It is interesting to note that while both states are correlated through eighth order in the MBPT, the singlet-triplet splitting is diminished with respect to our single-configuration HF splitting by |28.6 - 13.1| = 15.5 kcal/mol.

The final eighth-order MBPT splitting of -13.1 kcal/mol comes within 2.5 kcal/mol of the best extended basis set CI estimates of -10.6 kcal/mol.27,28 This can be considered excellent agreement given the modest (4s2p2d/2s) polarized basis function space over which the MBPT is defined in our calculations, since Bauschlicher and Shavitt got a full CI splitting of -12.2 kcal/mol when they employed a comparable, but slightly better, (4s2p2d/2s) basis set.29 Significantly, the extended polarized basis sets used in these other studies of 1 (e.g., (9s6p3d/4s2p), (6s4p2d/3s2p)29) also yielded HF-level 1A1-3B2 splittings whose magnitudes are about 4 kcal/mol smaller than our HF-level splitting of -28.6 kcal/mol (see Table IV). Thus, our results demonstrate that the MBPT method is capable of rendering adequate descriptions of both states of interest within our choice of basis.

The effectiveness of the two-configuration 1A1 and the one-configuration 1B2 valence descriptions in 1 suggests that considerations of valence electronic structure in 2 and 3 will provide good insights into the nature of the low-lying states of these unsaturated carbene as well. Even though the valence electronic structures of 2 and 3 are more complicated than the simple valence configurations A-C of 1, the descriptions of the lowest triplet states of these unsaturated carbene will be dominated by configurations analogous to A, while configurations analogous to B and C will contribute significantly to the lowest singlet state. The MBPT theory provides a convenient means for building a state description around such dominant configurations while automatically taking account of other significant configurations that manifest themselves in unsaturated carbene states. In what follows, we will make frequent reference to configurations that arise from various occupancies of the valence orbitals of 2 and 3. The relative importance of these configurations will be deduced from simple structure and bonding arguments.

**B. Vinylidene.** The dominant valence configurations of 2 corresponding to configurations A-C of 1 are D-F. To these configurations, we must also add another valence configuration, G.

Configuration G does not preserve the C-C π bond, but it is still favorable on the basis of orbital energetics (i.e., it can be viewed as a double excitation from the b2 π orbital, the highest occupied molecular orbital (HOMO) of the dominant singlet configuration C, to the lowest unoccupied molecular orbital (LUMO) of C, the b2 π orbital. Configurations D and F, on the other hand, involve excitations from the a1 σ HOMO-1 orbital of E). From configurations D-G for 2, which correspond to all possible valence configurations of the appropriate spin and spatial symmetry, we can write the state descriptions

\[ |3B2⟩_{HCCC} ≈ [\text{core}] a_1^2 b_2^2 \]  
\[ |1A1⟩_{HCCC} ≈ C_E[\text{core}] a_1^2 b_2^2 + C_F[\text{core}] a_1^2 b_2^2 + C_G[\text{core}] a_1^2 b_2^2 \]  

where \( |C_E|^2 + |C_F|^2 + |C_G|^2 \approx 1 \) in eq 4.

The core in carbene 2 consists of two filled carbon 1s atomic orbitals with a1 symmetry, a C-C σ bond with a1 symmetry, and two C-H bonding orbitals with a1 and b2 symmetry. The valence region is comprised of a π bonding orbital with b1 symmetry and, depending on the state or configuration, a π nonbonding orbital with b2 symmetry lying in the plane of the hydrogens, and a hybridized nonbonding σ orbital with a1 symmetry.

To understand the bonding and state ordering in 2, it is helpful to make comparisons between it and carbene 1. Directing our attention to the valence region, we note that carbene 2 differs from carbene 1 in three important ways. First, the π valence nonbonding orbital in 1 has b1 symmetry, but in 2 it has b2 symmetry and lies in the molecular plane. The possibility for a hyperconjugative interaction between the b2 C-H bond and the b2 nonbonding orbital must be considered in both the 1A1 and 1B2 states of 2. Second, the bonding b1 π orbital in the valence region of 2 has no analogue in carbene 1. As we have seen, double excitations from this orbital can give rise to another valence configuration contribution to the 1A1 state. Third, the orbitals localized on the carbene atoms of 1 and 2 are hybridized in different ways. For example, triplet state Mulliken population analysis shows that the four L shell (i.e., 2s2p2) carbene carbon electrons of 2 are hybridized as

\[ pp(sp)^2 \sim pb(1c\sigma\pi) pb(1b\pi) (sp)^2_{a1(c\sigma\pi)} (sp)^2_{a1(b\pi)} \]  

contrasting with the

\[ pp(sp)^2 + pb(1b\pi) (sp)^2_{a1(c\sigma\pi)} (sp)^2_{a1(b\pi)} \]  

hybridization found in triplet 1.26,29 Similarly, singlet 2 exhibits

\[ p(s^<p^<\alpha) (s^<p^<\alpha)^2 \sim pb(1c\sigma\pi) (s^<p^<\alpha)^2_{a1(c\sigma\pi)} \times (s^<p^<\alpha)^2_{a2(b\pi)} \]  

(carbene carbon hybridization),23 whereas the hybridization in singlet 1 is26,29

\[ (s^<p^<\alpha)^2 (s^<p^<\alpha)^2 \sim (s^<p^<\alpha)^2_{a1(c\sigma\pi)} \times (s^<p^<\alpha)^2_{a2(b\pi)} \]  

The singlet valence configurations B in 1 and E,G in 2 all arise from placement of the nonbonding electron pair in an a1 σ valence orbital. From eq 5-8, it is seen that the a1 σ nonbonding valence orbital in 2 has greater s character and hence lower orbital energy than the analogous orbital in 1 (e.g., ~0.46 hartree in singlet 2 vs. ~0.39 hartree in singlet 1). The greater s character of this valence orbital helps to make singlet orbital occupancy in 2 more favorable energetically.

In view of these differences between carbene 1 and 2, particularly carbene carbon orbital hybridization and the additional singlet configuration G, our prediction of a 1A1 ground state for 2 is not surprising. The ab initio calculations of Davis, Goddard, and Harding13 and Dykstra and Schaefer,14 as well as recent experimental data,3 also suggest singlet spin multiplicity for the ground state of 2. Early HF and semiempirical studies on this carbene predicted a singlet ground state as well.30-32 The correlation contributions of the 3B2 \( \rightarrow 2B2 \) and b2 \( \rightarrow b2^2 \) double excitation configurations F and G in 2 lower the energy of the 1A1 ground state, thereby increasing the maximum of the calculated 1A1-1B2 energy splitting relative to the splitting determined with single-configuration state descriptions (i.e., only configuration E for the 1A1 state, configuration D for the 1B2 state). Thus, we find that the 3B2 state.
of 2 lies 51.1 kcal/mol above the \(1A_1\) state in an eighth-order MBPT calculation where account is taken of the \(1A_1\) configurations F and G as well as all other double excitation configurations, but our HF-level single-configuration calculations place this excited triplet state only 15.7 kcal/mol above the \(1A_1\) state (see Table II). Since the \(1A_1\) state lies above the \(3B_1\) state in carbene I, the effect of the analogous \(a_1^2 \rightarrow b_2^2\) double excitation configuration C in this system is to diminish the magnitude of the \(1A_1\)-\(1B_1\) splitting relative to a HF-level splitting.

On the basis of semiemipirical MINDO/2 calculations, Bodor et al.\(^{31}\) predicted that the \(3B_2\) state of 2 lies 20.5 kcal/mol above the \(1A_1\) ground state, a result qualitatively confirmed in the work of Dykstra and Schaefer,\(^{14}\) who obtained a singlet–triplet splitting of 32.4 kcal/mol with their self-consistent electron pair (SCEP) method, which includes a modest treatment of electron correlation. Our MBPT results are in good agreement with the generalized valence bond (GVB) study of Davis, Goddard, and Harding,\(^{33}\) which, with careful account of correlation effects in both the \(1A_1\) and \(3B_2\) states, yielded a singlet–triplet splitting of 45.9 kcal/mol for 2 at slightly different state geometries from those we selected. The larger magnitude of the HF level singlet–triplet splitting in 2 reported by these authors (i.e., 31.0\(^{31}\) and 27.2 kcal/mol\(^{34}\) vs. our corresponding value of 15.7 kcal/mol) can be attributed to their use of a restricted Hartree–Fock (RHF) rather than a UHF triplet. The UHF necessarily yields a lower triplet energy than the RHF, thereby stabilizing the \(3B_2\) HF state relative to the \(1A_1\) HF state. Since the triplet is above the singlet in 2, the use of a UHF triplet gives rise to a smaller singlet–triplet splitting at the HF level.

C. Vinylidenecarbene. The dominant valence configurations of 3 analogous to A–C of 1 and D–F of 2 are shown in H–J. Carbene 3 also has an important group of valence configurations K–O arising from excitations out of the \(b_1\) and \(b_2\) bonding \(\pi\) orbitals of H–J. It is easily seen that configuration K, a \(b_2^2 \rightarrow b_2^2\) (i.e., HOMO \(\rightarrow\) LUMO) excitation of the singlet configuration I, is analogous to configuration G in carbene 2. However, the configurations L–O of 3 have no counterparts in either 1 or 2. The contributions of configurations H–O to the lowest singlet and triplet states of 3 are given by

\[
[3B_2]_{H\mbox{-}CCC} = C[H_{\mbox{core}} + a_1^2b_2^2b_2^2 + C_0_{\mbox{core}} + a_1^2b_2^2b_2^2 + C_{K_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{L_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{M_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{N_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{O_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{P_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{Q_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{R_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{S_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{T_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{U_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{V_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{W_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{X_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{Y_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{Z_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\alpha_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\beta_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\gamma_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\delta_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\epsilon_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\zeta_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\eta_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\theta_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\iota_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\kappa_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\lambda_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\mu_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\nu_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\xi_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\omicron_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\pi_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\rho_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\sigma_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\tau_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\upsilon_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\phi_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\chi_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\psi_{\mbox{core}}} a_1^2b_2^2b_2^2 + C_{\omega_{\mbox{core}}} a_1^2b_2^2b_2^2
\]
triplet (see Table IV).

It is interesting to note that Hehre et al. find the singlet state of OCS to lie above the triplet at the single-configuration HF level. As basis set quality is improved, however, the magnitude of the singlet–triplet splitting reported by these authors decreases from −25.6 kcal/mol (STO-3G basis) to only −1.5 kcal/mol (6-31G* basis including d-type polarization functions). These calculations illustrate that improvements to the basis set, particularly provisions for d-type polarization flexibility, bring about a significant preferential stabilization of the lowest singlet over the lowest triplet in a pattern also seen in theoretical studies of 1 and 2. Similarly, our preliminary HF test calculations on 3 employing a nonpolarized basis set also placed the triplet beneath the singlet by −4.8 kcal/mol. Since basis set augmentation and treatment of correlation, the two major ways to improve the quality of an ab initio theoretical study, both favor the singlet ground state, it is expected that any more complete study of 3 will confirm our predicted state ordering for this system. However, it should be kept in mind that our basis set for 3, which contains d functions only on the carbene carbon, is slightly biased against the 3B1 state. Our theoretical results for carbene 3 are summarized in Table III.

D. Trends and Predictions. Listed in Table IV are the A1 and 3B1,2 correlation energies, correlation energy differences, and uncorrelated and correlated energy splittings for carbones 1, 2, and 3. From the second and third columns of this table, we see that the magnitude of the correlation energy of both the A1 and 3B1,2 states increases as the complexity of the carbene species increases. In addition, the figures in the fourth column of the table indicate that the A1−3B1,2 energy difference increases with increasing carbene size (i.e., the correlation energy increase of the A1 state is proportionately larger than the increase of the 3B1,2 correlation energy as the carbene system itself becomes larger). This trend reflects the proliferation of singlet valence configurations as carbene size increases. At the HF level, however, the splitting between the A1 and 3B1,2 state diminishes in magnitude going from 2 to 3; the 3B1,2 HF state is consequently stabilized relative to the A1, HF state in the larger unsaturated carbones. Since the singlet state is more correlated than the triplet, a single-configuration approximation will always describe the triplet more accurately than the singlet. The latter trend would of course be expected to continue for extended unsaturated carbones such as 4 and 5, where increased nonbonding orbital delocalization would enhance the 3B1,2 HF state energy stabilization suggested by the trend in the fourth column of Table IV. Again, this follows from our observation that the number of configurations required to describe the singlet well increases more rapidly with carbene size than the number required for an equivalent treatment of the triplet. On the basis of these trends, we predict singlet ground states for carbones 4 and 5, since the strong correlation effects and π2−→ π2+ π2+ π2+ configuration delocalization influences in these systems would be expected to pull the A1 states beneath the triplet states, counteracting the "artificial" trend of 3B1,2 state stabilization at the HF level and the weak delocalization stabilization effect of the single nonbonding 3B1,2 π electron at the correlated level. Finally, it should be noted that given an extended valence π region, the lowest A2 states of 4 and 5, which can be thought to rise from b2π valence occupancy, may approach in energetic stability the corresponding 3B1,2 states of these species.

Our results also show that carbene carbon d functions must be present in basis sets for 2 and 3 to ensure accurate theoretical description of the low-lying states of these species. We find that the carbene carbon d functions preferentially stabilize the A1 states of 2 and 3. This effect is well documented in recent theoretical studies on carbene 1. For example, the erroneously large methylene (carbene 1) A1−3B1 splitting of −22.2 kcal/mol, computed by O'Neil, Schaefer, and Bender using the nonpolarized (4s2p/2s) basis together with multiconfiguration state descriptions,11 shows that even the multiconfiguration approach cannot completely compensate for a lack of basis set polarization flexibility. In the work of Meadows and Schaefer and Bauschlicher and Shavitt29 on 1, however, simple one- and two-configuration state descriptions, used in conjunction with excellent polarized HF-limit basis sets, gave the correct A1−3B1 splitting of −11 kcal/mol. Thus, it is not surprising to find that adequate theoretical description of the more correlated singlet states of unsaturated carbene systems requires a corresponding polarization flexibility in the basis. Without such flexibility, we find that even the state ordering in 3 is reversed at the HF level.

Conclusion

Our ab initio calculations predict a triplet ground state for methylene (1) and singlet ground states for both vinylidene (2) and vinylidenecarbene (3). The lowest excited state in each of these unsaturated carbene systems is found to have triplet spin multiplicity. Singlet–triplet energy splittings of −13.1, 51.1, and 48.7 kcal/mol were computed for 1, 2, and 3, respectively. The greater s character of the nonbonding a1 σ orbital on the terminal carbon and correlation effects arising from π2−→ π2+ π2+ π2+ configuration interaction of the valence electrons are identified as important factors in the stabilization of the singlet states of unsaturated carbones. Preferential stabilization of the singlet state in carbene 3 due to b1 σ orbital delocalization is discussed and its implications are noted for extended unsaturated carbones. Singlet ground states are predicted for the extended unsaturated carbones H2C::CCC: (4) and H2CCCC: (5) on the basis of singlet state correlation stabilization trends observed in carbones 2 and 3. The importance of carbon d-type polarization functions in unsaturated carbene basis sets is demonstrated. We find that this d function polarization flexibility preferentially stabilizes the singlet state, as in methylene.

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References and Notes

1. (a) University of Utah; (b) Battelle Columbus Laboratories.
A Molecular Orbital Study of Protonation. 5. Equilibrium Structures and Energies of Ions R₂CO⁺

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Abstract: Ab initio SCF calculations have been performed to determine the relative proton affinities of the carbonyl bases R₂CO and the structures of the ions R₂CO⁺, with R one of the isoelectronic saturated groups CH₃, NH₃, OH, and F. The predicted order of proton affinity with respect to R is NH₃ > CH₃ > OH > H > F, which is the same order predicted for the monosubstituted carbonyl bases RCO. Replacement of the hydrogen atom in RCHO by a second R group causes a further change in the proton affinity of the base in the same direction as observed upon substitution of the first R group, although the effect of two substituents is less than additive except in F₂CO. Protonation of carbonyl bases leads to an increase in the C-0 bond length and bond angles in the electron distribution upon protonation of the bases R₂CO are similar to the changes which occur upon protonation of the bases RCHO. From the computed results, a model for the protonation of carbonyl bases is proposed.

Introduction

In part 3 of this series,¹ the relative proton affinities of the bases RCHO were computed, and an analysis was made of substituent effects on proton affinities. In that study, the equilibrium structures of the relaxed ions RCHO⁺ were determined, and comparisons between the structures of these ions and the structures of the corresponding bases RCHO provided insight into the geometry changes due to protonation. That study has now been extended to the disubstituted carbonyls R₂CO, with R comprising the isoelectronic saturated groups CH₃, CH₂, OH, and F.


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