# The Importance of Including Dynamic Electron Correlation in ab Initio Calculations 

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A very useful simplifying assumption in chemistry is that the electrons in only a small number of orbitals are involved in most molecular processes. For example, the $\mathrm{N} \rightarrow \mathrm{V}$ transition in ethylene is usually described as involving the excitation of one electron from the $\pi$ to the $\pi^{*}$ molecular orbital. Similarly, the J ahn-Teller distortions ${ }^{1}$ of the radical ions of both aromatic and antiaromatic annulenes (e.g., benzene and cyclobutadiene) are attributed to the presence or absence of a single electron in a degenerate pair of $\pi$ molecular orbitals (MOs). ${ }^{2}$

Two electrons form the chemical bond in $\mathrm{H}_{2}$. Other localized chemical bonds, for example, the weak bond in the plane of the ring in 1,2-, 1,3-, and 1,4-dehydrobenzene ( $\mathbf{1}-\mathbf{3}$ ) are also commonly considered to involve just two electrons. The six $\pi$ electrons in 1-3 obviously do not participate in the weak in-plane bonds, because the sextet of $\pi$ electrons occupy MOs that are orthogonal to the $\sigma$ framework of the benzene ring.


1


2


3

Although more than just one or two electrons are involved in most chemical reactions, the number of electrons that participate is still a small fraction of the total number present in the reactants. For instance, as illustrated for the pericyclic reactions in Figure 1, the electrocyclic ring opening of cyclobutene is considered to involve just four electrons, two in the $\pi$ bond and two in the scissile $\sigma$ bond of the reactant. Six electrons, four in the two $\pi$ bonds of 1,5 -hexadiene and two in the $\mathrm{C}_{3}-\mathrm{C}_{4} \sigma$ bond, are depicted as participating in the Cope rearrangement. The electrons that

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Figure 1. Schematic depiction of three Woodward-H offmann allowed, pericyclic reactions: conrotatory electrocyclic ring opening of cyclobutene to 1,3-butadiene, [3,3]sigmatropic, chair Cope rearrangement of 1,5-hexadiene, and Diels-Alder cydoaddition of ethylene and 1,3-butadiene. Each arrow shows how one pair of "active" electrons is redistributed in each reaction.
are "active" in the Diels-Alder reaction between ethylene and butadiene are the six in the three $\pi$ bonds of the reactants.
It is tempting to attempt to transfer this conceptual simplicity to computational methodology. Indeed, semiempirical theories, such as Pariser-Parr-Pople (PPP), that deal only with the electrons in the $\pi \mathrm{MOs}$ of unsaturated hydrocarbons have been very successful in computing the UV spectra of these molecules. ${ }^{3}$ Therefore, one might expect that, in performing ab initio calculations on the $\pi \rightarrow \pi^{*}$ excitation in ethylene, focusing on just the two electrons in the $\pi$ orbitals would also be successful.

Similarly, since the presence of a single electron in a degenerate pair of $\pi \mathrm{MOs}$ is responsible for the J ahn-Teller distortion that is predicted for cyclobutadiene radical cation (CBD ${ }^{++}$), one might also anticipate that an independent particle model should satisfactorily predict whether a rectangle or a paralIelogram is the distorted geometry of lower energy. Therefore, Hartree-Fock (HF) calculations, either restricted (RHF) or unrestricted (UHF), would be expected to provide at least the qualitatively correct equilibrium geometry of CBD ${ }^{+}$.

HF calculations on the dehydrobenzenes are less likely to be successful, since the weak in-plane bonds in 1-3 are expected to have appreciable amounts of diradical character. At least two configurations are necessary to provide correlation between the pair of

[^0]electrons in a singlet diradical. ${ }^{4}$ Therefore, satisfactory wave functions for the dehydrobenzenes are likely to require two-configuration (TC)SCF calculations that correlate the pairs of electrons which form the weak in-plane bonds in 1-3.

Although HF calculations should, in principle, furnish a satisfactory description of the symmetryallowed ${ }^{5}$ pathways for the pericyclic reactions in Figure 1, the global potential surfaces for these reactions contain regions where diradical structures exist. ${ }^{6}$ Consequently, in order to calculate in an unbiased way the global potential surfaces for these reactions, multiconfigurational wave functions really are necessary.

A logical approach would be to provide correlation for all the electrons that are "active" in these reactions. Complete active space SCF (CASSCF) calculations correlaten "active" electrons ( $\mathrm{n}=4 \mathrm{in}$ cyclobutene ring opening and $\mathrm{n}=6$ in the Cope rearrangement and Diels-Alder reaction) by generating all possible configurations which distribute these electrons among n orbitals." CASSCF wave functions for the "active" electrons in pericydic reactions might be expected to yield computational results that would be at least qualitatively and, perhaps, even quantitatively correct.
In this Account we describe the results of calculations on the nature of the lowest energy singlet excitation in ethylene ${ }^{8}$ and in a much larger molecule, 1,2,4,5-tetramethylenebenzene. ${ }^{9}$ We review the computational work on molecular distortions in CBD ${ }^{+10}$ and in other radical ions. ${ }^{11-13}$ We discuss calculations on the Cope rearrangement, ${ }^{14-16}$ cyclobutene ring opening, ${ }^{17}$ and theDiels-Alder reaction. ${ }^{18}$ Finally, we describe computational investigations of the dehydrobenzenes, ${ }^{19}$ the results of which led to a redetermination of the experimental heats of formation of these molecules. ${ }^{20,21}$

The ab initio studies discussed in this Account have all shown that computational methodologies which provide correlation for only the "active" electrons give results that are not only quantitatively but, in many cases, also qualitatively incorrect. Only when dynamic

[^1]correlation (i.e., correlation between the active and inactive electrons) is included do ab initio cal culations correctly predict: (1) $\mathrm{N} \rightarrow \mathrm{V}$ in ethylene is a $\pi \rightarrow \pi^{*}$ valence transition, not an excitation to a Rydberg state. ${ }^{8}$ (2) Rectangular CBD ${ }^{+}$is lower in energy than planar rhomboidal CBD ${ }^{+} .{ }^{10}$ (3) There is only one $\mathrm{C}_{2 h}$ stationary point on the potential surface for the chair Cope rearrangement; and it is the "aromatic" transition state for a concerted reaction pathway. ${ }^{15,16}$

## Dynamic Electron Correlation in Ethylene

Ethylene is the prototypical molecule for the attempted separation of a many-electron system into two active electrons, with the remaining electrons treated as a passive core. Electronic states of ethylene can be formed by distributing the active el ectrons between the $\mathrm{p}-\pi$ atomic orbitals (AOs), $\mathrm{p}_{\mathrm{A}}$ and $\mathrm{p}_{\mathrm{B}}$, on the two carbon atoms. In valence bond (VB) theory a purely diradical, singlet wave function, $\Psi_{\mathrm{D}}$, can be written as

$$
\begin{equation*}
\Psi_{\mathrm{D}}=\left(\left|\mathrm{p}_{\mathrm{A}}{ }^{\alpha} \mathrm{p}_{\mathrm{B}}{ }^{\beta}\right\rangle-\left|\mathrm{p}_{\mathrm{A}}{ }^{\beta} \mathrm{p}_{\mathrm{B}}{ }^{\alpha}\right\rangle\right) / \sqrt{ } 2 \tag{1}
\end{equation*}
$$

where the symbol |> stands for a normalized Slater determinant, and $\alpha$ and $\beta$ are the two possible electron spins. In the wave function for the lowest triplet state, $\Psi_{T}$, one electron also occupies each AO, but the electron spins are parallel.

Neither $\Psi_{D}$ nor $\Psi_{T}$ contains ionic terms, corre sponding to the simultaneous occupancy of the same $\mathrm{p}-\pi$ AO by the two electrons. However, there are two totally ionic, singlet wave functions, which place both electrons on the same atom.

$$
\begin{align*}
& \Psi_{1}=\left(\left|\mathrm{p}_{\mathrm{A}}{ }^{\alpha} \mathrm{p}_{\mathrm{A}}^{\beta}\right\rangle+\left|\mathrm{p}_{\mathrm{B}}{ }^{\alpha} \mathrm{p}_{\mathrm{B}}{ }^{\beta}\right\rangle\right) / \sqrt{ } 2  \tag{2}\\
& \Psi_{\mathrm{V}}=\left(\left|\mathrm{p}_{\mathrm{A}}^{\alpha} \mathrm{p}_{\mathrm{A}}^{\beta}\right\rangle-\left|\mathrm{p}_{\mathrm{B}}{ }^{\alpha} \mathrm{p}_{\mathrm{B}}{ }^{\beta}\right\rangle\right) / \sqrt{ } 2 \tag{3}
\end{align*}
$$

$\Psi_{V}$ has $\mathrm{B}_{1 u}$ symmetry and is the wave function for the lower energy excited singlet state of ethylene. $\Psi_{1}$, like $\Psi_{D}$, has $A_{g}$ symmetry. The wave function, $\Psi_{N}$, for the ground state of ethylene, with energy $\mathrm{E}_{\mathrm{N}}$, contains both diradical and ionic terms; and it can be written as a linear combination of $\Psi_{D}$ and $\Psi_{1}$, with $C_{D}$ $>\mathrm{cl}$.

$$
\begin{equation*}
\Psi_{N}=c_{D} \Psi_{D}+c_{1} \Psi_{1} \tag{4}
\end{equation*}
$$

The singlet wave functions- $\Psi_{\mathrm{D}}, \Psi_{1}$, and $\Psi_{\mathrm{v}}$-for the two $\pi$ electrons in ethylene can also be described in molecular orbital (MO) theory. The bonding and antibonding $\pi$ MOs in ethylene can be written, respectively, as

$$
\begin{gather*}
\pi=\left(\mathrm{p}_{\mathrm{A}}+\mathrm{p}_{\mathrm{B}}\right) / \sqrt{ } 2  \tag{5}\\
\pi^{*}=\left(\mathrm{p}_{\mathrm{A}}-\mathrm{p}_{\mathrm{B}}\right) / \sqrt{ } 2 \tag{6}
\end{gather*}
$$

where, for the sake of simplicity, overlap has been neglected in normalizing the MOs. Using these MOs, three singlet configurations can be constructed. Expressed in terms of these three MO configurations, the
three singlet VB wave functions in eqs 1-3 can be written

$$
\begin{align*}
& \Psi_{\mathrm{D}}=\left(\left|\pi^{\alpha} \pi^{\beta}\right\rangle-\left|\pi^{* \alpha} \pi^{* \beta}\right\rangle\right) / \sqrt{ } 2  \tag{7}\\
& \Psi_{\mathrm{I}}=\left(\left|\pi^{\alpha} \pi^{\beta}\right\rangle+\left|\pi^{* \alpha} \pi^{* \beta}\right\rangle\right) / \sqrt{ } 2  \tag{8}\\
& \Psi_{\mathrm{V}}=\left(\left|\pi^{\alpha} \pi^{* \beta}\right\rangle-\left|\pi^{\beta} \pi^{* \alpha}\right\rangle\right) / \sqrt{ } 2 \tag{9}
\end{align*}
$$

In terms of MO configurations, the wave function for the ground state of ethylene thus becomes

$$
\begin{equation*}
\Psi_{\mathrm{N}}=\mathrm{c}_{\pi}\left|\pi^{\alpha} \pi^{\beta}\right\rangle-\mathrm{c}_{\pi} *\left|\pi^{* \alpha} \pi^{* \beta}\right\rangle \tag{10}
\end{equation*}
$$

Using eqs 8 and 9 and equating the VB and MO expressions for $\Psi_{N}$ in eqs 4 and 10 gives

$$
\begin{align*}
& \mathrm{C}_{\mathrm{D}}=\left(\mathrm{C}_{\pi}+\mathrm{C}_{\pi^{*}}\right) / \sqrt{ } 2  \tag{11}\\
& \mathrm{C}_{1}=\left(\mathrm{c}_{\pi}-\mathrm{C}_{\pi^{*}}\right) / \sqrt{ } 2
\end{align*}
$$

When, as is the case in Hartree-Fock theory, the wave function for the ground state consists of a single configuration ( $c_{\pi^{*}}=0$ in eq 10), eq 11 shows that the wave function for the ground state in eq 4 is an equal mixture of diradical and ionic terms. In contrast, when $\mathrm{c}_{\pi}=\mathrm{c}_{\pi^{*}}$, as would be the case in the transition state for rotation about the double bond in ethylene, where $\pi$ and $\pi^{*}$ are degenerate in energy, eq 11 shows that $\Psi_{N}$ is a pure diradical wavefunction and contains no ionic terms.

As indicated by eq 4, the actual wave function for the ground state of ethylene will contain a mixture of diradical and ionic terms, with the former dominant. The ratio of the coefficients, $C_{D}$ and $c_{1}$, depends on the energy difference, $\mathrm{E}_{1}-\mathrm{E}_{\mathrm{D}}$, between $\Psi_{1}$ and $\Psi_{\mathrm{D}}$ and on the size of the matrix element, $\left\langle\Psi_{\mathrm{I}}\right| \mathrm{H}\left|\Psi_{\mathrm{D}}\right\rangle$, that mixes them.

If the two $\mathrm{CH}_{2}$ groups in ethylene were separated by an infinite distance, $E_{v}$ and $E_{1}$ would be exactly equal and would differ from $E_{D}$ by IP - EA, the ionization potential minus electron affinity of each $\mathrm{CH}_{2}$ group. In the PPP model ${ }^{3}$ this energy difference is just the Coulomb repulsion energy between a pair of electrons in a $\mathrm{p}-\pi \mathrm{AO}$, which is symbolized as $\gamma \mathrm{cc}$. In PPP theory $\gamma \mathrm{cc}$ is set equal to the experimental value of 11 eV for IP - EA for the valence state of a free carbon atom. A better model for $\gamma_{\mathrm{cc}}$ in ethylene would be the experimental value of IP - EA for methyl radical, for which $I P-E A=9 \mathrm{eV}$.

Even at the HF level of ab initio theory, the $\sigma$ electrons in $\mathrm{CH}_{3}{ }^{-}, \mathrm{CH}_{3}{ }^{\circ}$, and $\mathrm{CH}_{3}{ }^{+}$will all have different wave functions, because of the difference in the number of electrons in the $p-\pi A O$. The different electrostatic polarizations of the $\sigma$ bonds in these three species are easily accommodated in separate ab initio calculations on $\mathrm{CH}_{3}{ }^{-}, \mathrm{CH}_{3}{ }^{\text {, }}$, and $\mathrm{CH}_{3}{ }^{+}$. However, in ab initio calculations on ethylene, the $\sigma$ polarizations in $\Psi_{।}$ and $\Psi_{V}$ are more difficult to handle, because they are dynamic (i.e., specific to each of the two $p-\pi$ AO occupancies in eqs 2 and 3), rather than static (i.e., based on the average $\pi$ charge distribution for the wave function in each equation).

In $\Psi_{1}$ and in $\Psi_{V}$, as in $\Psi_{\mathrm{D}}$, there is, on average, no net charge on either $\mathrm{CH}_{2}$ group of ethylene. Therefore, if $\Psi_{\mathrm{D}}{ }^{\text {core }}$ is the wave function for the core of $\sigma$ electrons that is appropriatefor the diradical wavefunction, $\Psi_{\mathrm{D}}$, for the two $\pi$ electrons, $\Psi_{D}{ }^{\text {core }}$, or something close to it, will also be the best average wave function for $\Psi_{1}$ and $\Psi_{V}$. However, the best total wave function for the V state of ethylene is not the HF wave function

$$
\begin{align*}
\Psi_{V}{ }^{\mathrm{HF}} \approx\left|\Psi_{\mathrm{D}}^{\text {core }} \Psi_{\mathrm{V}}\right\rangle=\left(\left|\Psi_{\mathrm{D}}^{\text {core }} \mathrm{p}_{\mathrm{A}}^{\alpha} \mathrm{p}_{\mathrm{A}}^{\beta}\right\rangle-\right. \\
\left.\left|\Psi_{\mathrm{D}}^{\text {core }} \mathrm{p}_{\mathrm{B}}^{\alpha} \mathrm{p}_{\mathrm{B}}^{\beta}\right\rangle\right) / \sqrt{ } 2 \tag{12}
\end{align*}
$$

The HF wave function does not allow the core of $\sigma$ electrons to become appropriately pol arized when both $\pi$ electrons are in $p_{A}$, leaving $p_{B}$ empty, and viceversa. Dynamic polarization requires a wave function in which dynamic correlation between the $\sigma$ and $\pi$ electrons is included.

Suppose that, when $p_{A}$ is doubly occupied and $p_{B}$ is empty, the appropriately polarized wave function for the $\sigma$ core is $\Psi_{\mathrm{A}^{-} \mathrm{B}^{+}}{ }^{\text {core }}$. Then an unnormalized wave function for the $V$ state of ethylene which includes dynamic correlation is

$$
\begin{equation*}
\Psi_{\mathrm{V}}{ }^{\text {dyn }}=\left|\Psi_{\mathrm{A}^{-} \mathrm{B}^{+}}{ }^{\text {core }} \mathrm{p}_{\mathrm{A}}^{\alpha} \mathrm{p}_{\mathrm{A}^{\beta}}\right\rangle-\left|\Psi_{\mathrm{A}^{+} \mathrm{B}^{-}}{ }^{\text {core }} \mathrm{p}_{\mathrm{B}}^{\alpha} \mathrm{p}_{\mathrm{B}}^{\beta}\right\rangle \tag{13}
\end{equation*}
$$

$\Psi_{\mathrm{D}}{ }^{\text {core }}$ in eq 12 is approximately the average of $\Psi_{\mathrm{A}^{-} \mathrm{B}^{+}}{ }^{\text {core }}$ and $\Psi_{\mathrm{A}^{+} \mathrm{B}^{-}}$core. Also, $\left(\Psi_{\mathrm{A}^{-} \mathrm{B}^{+}}{ }^{\text {core }}-\Psi_{\mathrm{A}^{+} \mathrm{B}^{-}}{ }^{\text {core }}\right) / 2$ can, to a first approximation, be written as a sum of single excitations, $\Sigma \Psi^{*}{ }^{\text {core }}$, from $\Psi_{\mathrm{D}}{ }^{\text {core }}$. Thus, the dynamically correlated wave function, $\Psi_{V}{ }^{\text {dyn }}$, in eq 13 can be obtained from the HF wave function, $\Psi_{V}{ }^{\mathrm{HF}}$, in eq 12 by adding to the latter the terms

$$
\begin{gather*}
\left.\left.\Psi_{V}{ }^{\text {dyn }}-\Psi_{V}{ }^{\mathrm{HF}}=\mid \sum \Psi^{*} \text { core }\left(\mathrm{p}_{\mathrm{A}}^{\alpha} \mathrm{p}_{\mathrm{A}}^{\beta}\right\rangle+\mathrm{p}_{\mathrm{B}}^{\alpha} \mathrm{p}_{\mathrm{B}}^{\beta}\right\rangle\right)= \\
\left.\left|\sum \Psi^{*}{ }^{\text {core }} \Psi_{\mathrm{I}}\right\rangle=\left|\sum \Psi^{*}{ }^{\text {core }}\left(\pi^{\alpha} \pi^{\beta}\right\rangle+\pi^{* \alpha} \pi^{* \beta}\right\rangle\right) \tag{14}
\end{gather*}
$$

Equation 14 shows that in ab initio MO theory provision of dynamic electron correlation for the V state of ethylene involves a sum of double excitations of one core electron and one active electron from the HF wave function for $\Psi_{V}$. Since $\Psi_{V}$ is ${ }^{1} \mathrm{~B}_{1 u}$, and $\Psi_{1}$ is ${ }^{1} \mathrm{~A}_{\mathrm{g}}$, only those terms in $\sum \Psi^{*}$ core that have ${ }^{1} \mathrm{~B}_{1 u}$ symmetry contribute to eq 14.

A similar expression can be easily derived for the terms that must be added in order to provide dynamic correlation for the ionic component, $\Psi_{1}$, of the ground state wave function, $\Psi_{N}$, for ethylene.

$$
\begin{align*}
& \Psi_{1}^{d y n}-\Psi_{I}^{\mathrm{dF}}=\left|\sum \Psi^{*}{ }^{\text {core }} \Psi_{\mathrm{V}}\right\rangle= \\
&\left.\left|\sum \Psi^{\text {core }}\left(\pi^{\alpha} \pi^{* \beta}\right\rangle-\pi^{\beta} \pi^{* \alpha}\right\rangle\right) \tag{15}
\end{align*}
$$

As with $\Psi_{V}$, provision of dynamic correlation between active and inactive electrons in $\Psi_{\text {I }}$ requires the simultaneous excitation of one electron of each type from the HF wave function. This is generally the case.

Including dynamic electron correlation for $\Psi_{\mid}$would be expected to increase its contribution to $\Psi_{N}$, the wave function for the ground state of ethylene. In fact, the ratio $\mathrm{C}_{\pi} * \mathrm{c}_{\pi}$ decreases from 0.21 in a CASSCF calculation with a very large basis set to 0.16 in a multireference (MR) configuration interaction (CI) calculation with the same basis set. ${ }^{8}$ This change corresponds to an increase in the ratio $c_{I} / c_{D}$ from 0.65 to 0.72 . The dynamic correlation provided in the

| state | calculation ${ }^{\text {b }}$ | $\Delta \mathrm{E}$ | $\left\langle x^{2}\right\rangle$ |
| :---: | :---: | :---: | :---: |
| T | CASSCF | 4.27 | 11.7 |
| v | CASSCF | 8.05 | 43.4 |
| v | CASSCF (with N MOs) | 12.33 | 10.9 |
| T | $\pi \mathrm{Cl}$ | 4.34 | 11.7 |
| v | $\pi \mathrm{Cl}$ | 8.09 | 40.1 |
| v | $\pi \mathrm{Cl}$ (with N MOs) | 8.73 | 26.3 |
| T | ASSEFAPC | 4.41 | 11.7 |
| $v$ | ASSEFAPC | 8.13 | 18.5 |
| v | ASSEFAPC (with N MOs) | 8.06 | 18.8 |

MRCI calculation increases the occupancy of the $\pi$ MO, at the expense of $\pi^{*}$, by stabilizing the ionic terms in the $\left|\pi^{2}\right\rangle$ configuration.

## Calculations on Excited States

Ethylene. Dynamic electron correlation should be even more important in $\Psi_{V}$ than $\Psi_{N}$, since the wave function for the $\pi$ electrons in the first excited singlet state of ethylene is purely ionic. The results of recent ab initio cal culations on ethylene with a very extended basis set show that whether the V state is computed to be valence or Rydberg depends on whether dynamic electron correlation is included in the wave function for this state. 8,22 Table 1 gives the exditation energies, computed with several different types of wave functions, including wave functions in which the MOs from CASSCF calculations on the ground ( N ) state were used to construct the wave function for the V excited state.

The CASSCF MOs for V are very different from those for N and $\mathrm{T} .{ }^{23}$ This is shown by the values of $\left\langle x^{2}\right\rangle$ for these states in Table 1. The quantity $\left\langle x^{2}\right\rangle$ for each state gives the spatial extent of the wave function out of the molecular plane and is thus indicative of the size of the $\pi$ MOs in each state. In the CASSCF wave function, $\Psi_{\mathrm{N}}$, for the ground state of ethylene, $\pi^{*}$ is used to correl ate the two electrons in the bonding $\pi$ MO. Therefore, $\pi^{*}$ must overlap effectively with $\pi$. Consequently, these two MOs have similar sizes, giving $\left\langle x^{2}\right\rangle=10.9$ bohr $^{2}$ for $\Psi_{\mathrm{N}}$. The CASSCF triplet wave function, $\Psi_{T},{ }^{23}$ has about the same value of $\left\langle x^{2}\right\rangle$ as $\Psi_{N}$, because the $\pi$ and $\pi^{*} \mathrm{MOs}$ in $\Psi_{\mathrm{T}}$ have about the same size as the $\pi$ and $\pi^{*}$ MOs in $\Psi_{N} .^{24}$

However, in the CASSCF wave function for the excited singlet state, $\Psi{ }_{v},{ }^{23}$ the value of $\left\langle x^{2}\right\rangle$ is about 4 times larger than in either $\Psi_{N}$ or $\Psi_{T}$. The reason is that, as shown in eq 3, the two $\pi$ electrons in $\Psi_{v}$ are

[^2]always on the same atom. Therefore, if the sizes of the AOs in the $\pi$ and $\pi^{*}$ MOs were similar, the two $\pi$ electrons would have a very high probability of simultaneously being in the same region of space and, thus, would have a very high Coulomb repulsion energy. The diffuse, Rydberg-like $\pi^{*}$ MO of $\Psi_{V}$ keeps the electron in this MO confined to a region of space that is quite different from that where the electron in the much more contracted $\pi \mathrm{MO}$ is most likely to be found. Confining electrons of opposite spin to different regions of space minimizes their Coulombic repulsion energy. ${ }^{1,25}$
The results in Table 1 show that the energy of the $\mathrm{N} \rightarrow \mathrm{V}$ transition in ethylene depends on the type of MOs used to construct the V excited state. Most dramatic is the finding that using the MOs from the CASSCF (two electrons in two orbitals) calculation on $\Psi_{N}$ to construct $\Psi_{V}$ gives a $\mathrm{N} \rightarrow$ V CASSCF excitation energy that is 4.28 eV higher than when a CASSCF calculation is actually performed on the V state. ${ }^{23}$

When full Cl is carried out in the $\pi$ space, the $\pi$ wave function for the V state would be independent of whether N or V CASSCF MOs were used, provided that the CASSCF MOs for the $\sigma$ cores of the two states were the same. The results in Table 1 show that the $\sigma$ MOs from these two calculations are obviously quite different. The value of $\left\langle x^{2}\right\rangle$ for the V state is highly dependent on whether the $\sigma$ core wave function from the CASSCF calculation on V or N is used in the $\pi \mathrm{Cl}$ calculation.
Dynamic correlation between the $\sigma$ core and the two $\pi$ electrons can be included by adding single $\sigma$ excitations to all the $\pi$ configurations. The resulting ASSEFAPC (all single $\sigma$ excitations from all $\pi$ configurations) Cl wave functions for N and V give energies and values of $\left\langle x^{2}\right\rangle$ that are nearly independent of whether the MOs from the CASSCF calculations on the N or the V state are used. Because dynamic correl ation stabilizes ionic terms in the wave function for $\Psi_{v}, \pi^{*}$ has more nearly the same size as $\pi$ in the ASSEFAPC wave function than in the CASSCF or $\pi$ CI wave functions.
Using the appropriate CASSCF MOs for T and V , the vertical excitation energies for these two states are within 0.2 eV of experiment ${ }^{22 \mathrm{~h}}$ at all levels of theory in Table 1. However, the predicted nature of the V excited state depends critically on the level of theory used. At the CASSCF and $\pi \mathrm{Cl}$ levels the values of $\left\langle x^{2}\right\rangle$ in Table 1 indicate that the $V$ state of ethylene is a Rydberg state. In contrast, with inclusion of dynamic electron correlation at the ASSEF APC level, the values of $\left\langle x^{2}\right\rangle=18.5-18.8$ for the V state indicate that, although the $\pi^{*} \mathrm{MO}$ in it is more diffuse than the $\pi^{*} \mathrm{MO}$ in the T state, the V state of ethylene is not a Rydberg state.

The $\mathbf{S}_{\mathbf{0}} \boldsymbol{\rightarrow} \mathbf{S}_{\mathbf{1}}$ Excitation in TMB (4). 1,2,4,5Tetramethylenebenzene (TMB) is a diradical whose nonbonding (NB) MOs can be chosen so that they are essentially those for two pentadienyl radicals and, thus, span disjoint sets of atoms. Consequently, both qualitative considerations ${ }^{25}$ and ab initio calculations ${ }^{9,26}$ predict that TMB will violate Hund's rule ${ }^{27}$ by having a singlet ground state. However, when this

[^3]diradical was first prepared, several observations led Roth and co-workers to identify the triplet as the ground state. ${ }^{28}$


One of these observations was that the UV-vis spectrum of $\mathbf{4}$ consists of a weak absorption at long wavelengths, followed by a much more intense absorption. Both $\pi \mathrm{Cl}^{26}$ and CASSCF calculations ${ }^{9}$ predict that the lowest energy vertical excitation of singlet TMB is an allowed transition, followed by a forbidden excitation at higher energy. However, the observed spectrum fits the pattern predicted for the triplet by $\pi \mathrm{CI}$ and CASSCF calculations, a weak forbidden absorption followed by a strong allowed absorption.
The excitation energies of singlet and triplet TMB have recently been recal culated at the CASPT2N level of theory. ${ }^{9}$ CASPT2N uses second-order MöllerPlesset (MP2) perturbation theory to provide dynamic electron correlation for a CASSCF wave function. ${ }^{29} \mathrm{~A}$ closely related CASMP2 method for adding dynamic correlation to a CASSCF wave function has also been published by one of us. ${ }^{30}$

At the CASPT2N level of theory the vertical excitation $1^{1} \mathrm{~A}_{g} \rightarrow 2^{1} \mathrm{~A}_{g}$, which is forbidden, falls slightly bel ow $1^{1} \mathrm{~A}_{g} \rightarrow{ }^{1} \mathrm{~B}_{3 u}$, which is allowed. Moreover, upon optimization of the geometries of these two excited states, the greater relaxation energy computed for $2^{1} \mathrm{~A}_{g}$ results in the adiabatic excitation energy for this state being about 0.2 eV lower than that for ${ }^{1} \mathrm{~B}_{3 \mathrm{u}}$. The UVvis spectrum for singlet TMB, computed by CASPT2N, ${ }^{9}$ provides not only a qualitative but also a quantitative fit to the UV-vis spectrum that was observed by Roth and co-workers. ${ }^{28}$

The UV-vis spectrum for the triplet, computed at the CASPT2N level, ${ }^{9}$ does not fit the observed spectrum. The CASPT2N calculations thus provide strong evidence that the UV-vis spectrum of TMB belongs to the singlet and not to the triplet. Berson and coworkers ${ }^{31}$ have obtained other types of experimental evidence that, as predicted ${ }^{9,25,26}$ the singlet is, in fact, the ground state of TMB. ${ }^{27}$

## Radical Cations

Cyclobutadiene Radical Cation (CBD ${ }^{+}$). As discussed in the previous section, when the $\pi$ wave function for an excited state of a neutral molecule contains ionic terms, dynamic electron correlation between $\pi$ and $\sigma$ electrons is very important. Therefore, it is not surprising that this type of correlation is al so important in conjugated molecules with charged $\pi$ systems. A good example is provided by CBD ${ }^{\cdot+} .^{10}$
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Figure 2. Rectangular ( $\mathrm{b}_{1 \mathrm{~g}}$ ) and rhomboidal ( $\mathrm{b}_{2 \mathrm{~g}}$ ) distortions of $\mathrm{D}_{4 h}$ CBD•+. Only the top lobes of the $\mathrm{p} A O s$ in the $\pi \mathrm{MO}$ that is singly occupied in the ground state at each distorted geometry are drawn. The phase change in the electronic wave function that is necessarily encountered upon completing one pseudorotation circuit about the $\mathrm{D}_{4 \mathrm{~h}}$ geometry (Longuet-Higgins, H. C. Proc. R. Soc. London, Ser. A 1975, 344, 147) is shown as occurring between the $\mathrm{b}_{2 g}$ distorted geometry at the bottom of the figure and the $b_{1 g}$ distorted geometry at the left.

As shown in Figure 2, square ( $\mathrm{D}_{4 \mathrm{~h}}$ ) CBD ${ }^{++}$can undergo two different types of J ahn-Teller distortions: $\mathrm{b}_{1 g}$, which takes the molecule to one of two rectangular geometries, and $\mathrm{b}_{2 \text { g, }}$, which takes it to one of two rhomboidal geometries. The radical cation can pseudorotate from one of these distorted geometries to another, but group theory cannot predict which geometry will be a minimum and which a transition state for pseudorotation.

One might expect that distortion of the $D_{4 h}$ bond lengths would provide more stabilization than distortion of the $\mathrm{D}_{4 \mathrm{~h}}$ bond angles, and our ab initio calculations confirmed that this is, indeed, the case. ${ }^{10}$ The $\mathrm{b}_{1 g}$ distortion lowers the energy by $7.0 \mathrm{kcal} / \mathrm{mol}$, whereas the energy lowering associated with the $b_{2 g}$ distortion is only $1.0 \mathrm{kcal} / \mathrm{mol}$. Nevertheless, at many levels of theory, the optimized rhomboidal geometry is lower in energy than the optimized rectangular geometry.

The reconciliation of these two, apparently contradictory, statements is that the energy of the square molecule depends on the type of degenerate MO that is singly occupied. As shown in Figure 2, a rectangular ( $\mathrm{b}_{19}$ ) distortion requires that the singly occupied NBMO be chosen to span all four atoms, whereas a rhomboidal ( $\mathrm{b}_{2 \mathrm{~g}}$ ) distortion requires that the singly occupied NBMO be confined to two carbons that are diagonally across the ring from each other. Although the computed energy of the square molecule should be independent of which type of NBMO is chosen, this does not mean that at approximate levels of theory the computed energy actually exhibits this independence.
At the UHF level the more localized choice for the form of the NBMO that is singly occupied gives an energy for square CBD ${ }^{++}$which is lower by $20.8 \mathrm{kcal} /$ mol than the less localized choice for this NBMO. ${ }^{10}$ Without explicit inclusion of electron correlation, localization is favorable, since the other $\pi$ and $\sigma$ electrons can adjust their wave functions to capitalize on the localization of spin and charge.

At the ROHF level the dependence of the energy of the square molecule on the type of degenerate NBMO chosen is smaller than at the UHF level, amounting to $7.4 \mathrm{kcal} / \mathrm{mol}$. With inclusion of correlation between the $\pi$ electrons via full Cl in the $\pi$ space, this difference drops to $4.8 \mathrm{kcal} / \mathrm{mol}$. The existence of this
difference, even when the $\pi$ electrons are fully correlated, indicates the importance of providing dynamic correlation between the $\pi$ and $\sigma$ electrons. When this was done by including all single $\sigma$ excitations from all $\pi$ configurations (ASSEFAPC), the dependence of the energy of the square molecule on the form chosen for the singly occupied NBMO decreased to $0.6 \mathrm{kcal} / \mathrm{mol}$, with the less localized NBMO now giving the lower energy.

The symmetry requirement that the energy of square CBD*+ should be independent of the type of degenerate NBMO that is occupied provides a convenient test for the quality of different types of wave functions. Testing for artifactual symmetry breaking ${ }^{32}$ in this fashion indicates whether the amount of dynamic electron correlation that has been included in the wave function is sufficient to allow the accurate calculation of the energy difference between rectangularly and rhomboidally distorted CBD•+. 33

Cyclooctatetraene Radical Anion (COT* ${ }^{-}$). In COT•- three electrons occupy a pair of degenerate $\pi$ MOs at the $D_{8 h}$ geometry of the planar radical anion. As in CBD*+, J ahn-Teller distortions of either the bond lengths or the bond angles of COT•- are possible, with one type of distortion leading to the energy minima along the pathway for pseudorotation and the other leading to the transition states connecting them. Also as in CBD**, calculations on COT•- at the geometry of highest symmetry with two different types of NBMOs provide a test for artifactual symmetry breaking.

Because the $\pi$ system of COT•- is larger than that of CBD**, at any level of theory the energies computed with the two different types of MOs differ less at the $D_{8 h}$ geometry of the former than they do at the $D_{4 h}$ geometry of the latter. ${ }^{11}$ As expected, ASSE FAPC CI calculations, which provide dynamic correlation between the $\pi$ and $\sigma$ electrons, show very little artifactual symmetry breaking. Moreover, ASSEFAPC CI calculations at the bond-alternated, equilibrium, $\mathrm{D}_{4 \mathrm{~h}}$ geometry of COT•- give a hyperfine coupling constant of $\mathrm{a}_{\mathrm{H}}=-3.21 \mathrm{G}$, which is in exact agreement with experiment. ${ }^{34}$

Trimethylenemethane Radical Cation (TMM ${ }^{+}$). Testing for artifactual symmetry breaking in a molecule, such as TMM ${ }^{++}$, which has a 3 -fold axis of symmetry, is slightly different than in molecules, such as CBD** ${ }^{+}$, with 4 -fold symmetry axes. ${ }^{32}$ Neither of the degenerate $\mathrm{e}_{x}^{\prime \prime}$ and $\mathrm{e}_{y}^{\prime \prime} \pi$ NBMOs in TMM•+, shown in Figure 3, individually has $\mathrm{D}_{3 \mathrm{~h}}$ symmetry; each has only $\mathrm{C}_{2 v}$ symmetry. One NBMO belongs to the $\mathrm{a}_{2}$ representation of $\mathrm{C}_{2 v}$; and, when it is singly occupied, the resulting wave function is ${ }^{2} \mathrm{~A}_{2}$. The other NBMO belongs to the $b_{1}$ representation and gives rise to a ${ }^{2} \mathrm{~B}_{1}$ state.

The nonuniform distribution of charge and spin in the NBMO of each of these two states affects the $\sigma$

[^4]

$e_{y}{ }^{\prime \prime}\left(b_{1}\right)$
Figure 3. Degenerate $\mathrm{e}^{\prime \prime}$ MOs in $\mathrm{D}_{3 \mathrm{~h}} \mathrm{TMM}^{++}$. Only the top lobes of the p AOs in each $\pi \mathrm{MO}$ are shown. The $\mathrm{C}_{2 v}$ symmetry designation of each MO is given in parentheses.
and $\pi$ electrons in planar TMM++ in such a way that both states distort away from a $\mathrm{D}_{3 \mathrm{~h}}$ geometry, as predicted by the J ahn-Teller theorem. ${ }^{1}{ }^{2} \mathrm{~A}_{2}$ has one short $\mathrm{C}-\mathrm{C}$ bond and two longer ones, whereas the opposite is the case for ${ }^{2} \mathrm{~B}_{1}$. One of the distorted geometries corresponds to the three minima along the pathway for pseudorotation in planar TMM••+ and the other to the three transition states that connect the minima. Ab initio calculations were performed to identify which of the two $\mathrm{C}_{2 v}$ geometries corresponds to which of these two types of stationary points and to compute the energy difference between them. ${ }^{12}$

UHF calculations placed the ${ }^{2} \mathrm{~A}_{2}$ optimized geometry $6.6 \mathrm{kcal} / \mathrm{mol}$ below that of ${ }^{2} \mathrm{~B}_{1}$. In contrast, UMP2, ROHF , and SD-CI calculations all found ${ }^{2} \mathrm{~B}_{1}$ to be the lower of the two by, respectively, $5.5,8.3$, and $2.7 \mathrm{kcal} /$ mol. However, when each of these calculations was performed at a $D_{3 h}$ geometry, the energy differences found between ${ }^{2} \mathrm{~A}_{2}$ and ${ }^{2} \mathrm{~B}_{1}$ were about the same size as at the optimized $C_{2 v}$ geometries for these two states. Of course, pure ${ }^{2} E_{x}{ }^{\prime \prime}$ and ${ }^{2} E_{y}$ " states would have had identical energies at $D_{3 h}$ geometries.

We had previously used group theory to investigate for the $(\mathrm{NH})_{3}{ }^{2+}$ diradical what types of dynamic correlation between the $\sigma$ and $\pi$ electrons must be included in order to obtain pure ${ }^{1} \mathrm{E}_{\mathrm{x}}$ and ${ }^{1} \mathrm{E}_{\mathrm{y}}$ ' states at $D_{3 n}$ geometries. ${ }^{35}$ We found that, in order to obtain wave functions which, even to first order, are pure ${ }^{1} E_{x}{ }^{\prime}$ and ${ }^{1} \mathrm{E}_{\mathrm{y}}{ }^{\prime}$, all single $\sigma$ excitations from all $\pi$ configurations (ASSEFAPC) must be included. The same type of analysis shows that the dynamic electron correlation provided by ASSE FAPC CI should also give wave functions that are, at least to first order, pure ${ }^{2} E_{x}{ }^{\prime \prime}$ and ${ }^{2} \mathrm{E}_{\mathrm{y}}$ " states of $\mathrm{D}_{3 \mathrm{~h}} \mathrm{TMM}{ }^{++}$.

Indeed, ASSEFAPC CI at $\mathrm{D}_{3 n}$ geometries of TMM•+ gave ${ }^{2} \mathrm{~A}_{2}$ and ${ }^{2} \mathrm{~B}_{1}$ wave functions that differed in energy only by $0.1 \mathrm{kcal} / \mathrm{mol}$. When the J ahn-Teller distorted, $\mathrm{C}_{2 v}$ geometry of each state was optimized at the ASSEFAPC CI level, the two geometries were computed to have the same energy to within $0.1 \mathrm{kcal} /$ mol. Thus, with proper inclusion of dynamic electron correlation, ab initio calculations predict that pseudorotation in TMM ${ }^{++}$occurs without any appreciable energy barrier. ${ }^{12}$ This prediction is qualitatively different from that which would have been made on the basis of the results of UHF , UMP2, ROHF , or SD-CI calculations.

Trimethylene Radical Cation (TM• ${ }^{+}$) and Related Radicals. Like planar TMM ${ }^{\bullet+}$, which might be formed upon ring opening of methylenecyclopropane radical cation, the $(0,0)$ geometry of $\mathrm{TM}^{\bullet+}(\mathbf{6})$ might be formed by ring opening of cyclopropane radical

[^5]cation (5). We have performed calculations not only on the former reaction ${ }^{12}$ but also on the latter. ${ }^{13}$


J ust as symmetry considerations allow one to test the adequacy of wave functions for CBD** and TMM•+, symmetry also allows one to test the adequacy of wave functions for ( 0,0 )-TM•+ (6). One can thus determine whether the symmetry breaking calculated for this radical cation at the UHF, ROHF , and CASSCF levels of theory ${ }^{13}$ is real or artifactual. ${ }^{32}$

In ( 0,0 )-TM ${ }^{++}$the symmetry principle that can be used is that the wave function for a nondegenerate state must have the same symmetry as the mol ecular geometry. If this is found not to be the case for a particular type of wave function, both the symmetry breaking in the wave function and the molecular distortion to lower symmetry that necessarily accompanies it are clearly artifactual.

In the case of $(0,0)-\mathrm{TM} \cdot+$, cal culations performed at $\mathrm{C}_{2 v}$ geometries with UHF, ROHF , and CASSCF wave functions showed the symmetry breaking that they exhibited to be artifactual. ${ }^{13}$ In each case a wave function with only $\mathrm{C}_{s}$ symmetry was found to be lower in energy than a wave function on which $\mathrm{C}_{2 v}$ symmetry was imposed.

None of these wave functions contains dynamic correlation between the electrons in the $\pi \mathrm{MOs}$ and those in the $\sigma$ MOs. However, when this type of dynamic correlation was provided by ASSEFAPC CI calculations, artifactual symmetry breaking of the wave function was not observed. Moreover, at this level of theory the optimized $\mathrm{C}_{2 v}$ geometry of $(0,0)$ TM ${ }^{++}$was found to be stable toward lengthening one $\mathrm{C}-\mathrm{C}$ bond and shortening the other. Thus, when dynamic electron correlation is provided at the ASSEFAPC level of theory, artifactual symmetry breaking in the wave function disappears, and the radical is stable toward real breaking of the molecular symmetry.

Providing dynamic electron correlation between $\sigma$ and $\pi$ electrons through ASSEFAPC CI calculations has also proven important for investigating molecular distortions from $\mathrm{C}_{2 v}$ to $\mathrm{C}_{s}$ symmetry in other radicals with three $\pi$ electrons. These include two more radical cations, $\mathrm{O}\left(\mathrm{CH}_{2}\right)_{2}{ }^{\bullet+36}$ and $\mathrm{CF}_{2}{ }^{\bullet+},{ }^{37}$ and neutral formyloxyl radical. 36,38 In each case, without adequate provision of dynamic electron correlation, artifactual symmetry breaking was encountered; and this obscured whether each radical actually preferred a $\mathrm{C}_{2 v}$ or a $\mathrm{C}_{\mathrm{s}}$ geometry.

## Potential Energy Surfaces for Pericyclic Reactions

Cope Rearrangement. It is clear why dynamic electron correlation might be important for obtaining qualitatively meaningful results in excited states that

[^6]

Figure 4. Nature of the $C_{2 n}$ species on the PES for the chair Cope rearrangement as the distance, R , between $\mathrm{C}_{1}$ and $\mathrm{C}_{6}$ (and $\mathrm{C}_{3}$ and $\mathrm{C}_{4}$ ) decreases from the top to the bottom of the figure. The leading terms in the wave functions for the two singlet diradical extremes, two allyls and cyclohexane-1,4-diyl, are given.
have highly ionic $\pi$ wave functions or in molecules with charged or very polar $\pi$ systems. However, one might not expect inclusion of dynamic electron correlation to be necessary for calculating the correct potential energy surface for a simple, WoodwardH offmann allowed, ${ }^{5}$ pericyclic reaction, ${ }^{6}$ such as the Cope rearrangement. Nevertheless, recent calculations in both of the authors' research groups have demonstrated clearly that calculations beyond the CASSCF level are necessary to obtain even a qualitatively correct potential energy surface (PES) for this reaction. ${ }^{15,16}$

All calculations agree that the lowest energy pathway for the Cope rearrangement of 1,5-hexadiene passes through a $\mathrm{C}_{2 \mathrm{~h}}$ geometry, ${ }^{6}$ and discussion of the reaction has focused on the electronic structure and energy of this species and whether it is a minimum or saddle point on the potential energy surface. As shown in Figure 4, at long $\mathrm{C}_{1}-\mathrm{C}_{6}$ (and $\mathrm{C}_{3}-\mathrm{C}_{4}$ ) distances, $R$, the $C_{2 h}$ structure separates into two allyl radicals, whose energy is known experimentally to be about 60 $\mathrm{kcal} / \mathrm{mol}$ above that of 1,5-hexadiene. ${ }^{39}$ The singlet wave function for this radical pair consists of the two dominant configurations that are given in Figure 4.

At a $\mathrm{C}_{1}-\mathrm{C}_{6}$ distance, R , that is close to a normal $\mathrm{C}-\mathrm{C}$ single bond length, the $\mathrm{C}_{2 h}$ structure is a chair diradical, cyclohexane-1,4-diyl. The energy of this diradical is not known experimentally, but thermochemical estimates place it some $40 \mathrm{kcal} / \mathrm{mol}$ above 1,5-hexadiene. ${ }^{40}$ As shown in Figure 4, although the first configuration in the wave function for this diradical is the same as that for two, singlet-coupled, allyl radicals, the second configuration is not.

The actual transition state for the chair Cope rearrangement has an enthalpy that is $33.5 \mathrm{kcal} / \mathrm{mol}$ greater than that of 1,5-hexadiene. ${ }^{41}$ The difference between the enthal py of the transition state and that of each of the two diradical extremes in Figure 4 is the amount by which concerted bond making and bond breaking make the energy of the transition statelower

[^7]than each of the two hypothetical transition states in which these two processes are completely decoupled.
A CASSCF wave function for the Cope rearrangement would be expected to have as its three most important configurations
\[

$$
\begin{array}{r}
\Psi=\mathrm{c}_{1}\left|\ldots 7 \mathrm{a}_{\mathrm{g}}^{2} 5 \mathrm{a}_{\mathrm{u}}^{2} 7 \mathrm{~b}_{\mathrm{u}}^{2}\right\rangle-\mathrm{c}_{2}\left|\ldots 7 \mathrm{a}_{\mathrm{g}}^{2} 5 \mathrm{a}_{\mathrm{u}}^{2} 8 \mathrm{a}_{\mathrm{g}}^{2}\right\rangle- \\
\mathrm{c}_{3}\left|\ldots 7 \mathrm{a}_{\mathrm{g}}^{2} 5 \mathrm{~b}_{\mathrm{g}}^{2} 7 \mathrm{~b}_{\mathrm{u}}^{2}\right\rangle-\ldots \tag{16}
\end{array}
$$
\]

Such a wave function has the flexibility to describe both of the diradicals shown in Figure 4, as well as the chair transition state, whose electronic structure is expected to be intermediate betweeen them.

CASSCF/6-31G* calculations on the chair Cope rearrangement find two $\mathrm{C}_{2 \mathrm{~h}}$ stationary points: cyclo-hexane-1,4-diyl, which is a stable intermediate with a short (1.641 $\AA$ ) $\mathrm{C}_{1}-\mathrm{C}_{6}$ bond length, and an "aromatic" transition state with a long ( $2.189 \AA$ ) $\mathrm{C}_{1}-\mathrm{C}_{6}$ bond length. ${ }^{14}$ The enthalpies of both are calculated to be more than $10 \mathrm{kcal} / \mathrm{mol}$ too high, compared to the experimental value of $\Delta \mathrm{H}^{\ddagger}=33.5 \mathrm{kcal} / \mathrm{mol} .{ }^{41}$ At the diyl geometry, $c_{2} / c_{1}=0.63$, while, at the aromatic transition state, the third configuration in eq 20 becomes slightly more important than the second, and $\mathrm{C}_{3} / \mathrm{C}_{1}=0.18$. In $\mathrm{C}_{2 \mathrm{~h}}$ symmetry the two stationary points are bond-stretch isomers, separated by a small energy barrier which reaches its maximum height at $R=1.957 \AA$. As would be expected from the discussion given above for ethylene, CASSCF underestimates the importance of the ionic terms in the wave functions for the active electrons. Lack of correlation between the 6 active and 28 inactive valence electrons effectively places the ionic terms too high in energy. Inclusion of this correlation by CASPT2N ${ }^{15}$ or CASMP2 ${ }^{16}$ reduces the importance of the second configuration in eq 16, which removes ionic terms from the first configuration at the geometry of cyclohexane-1,4-diyl. ${ }^{42}$

The CASPT2N and CASMP2 PESs each have a single $\mathrm{C}_{2 h}$ stationary point at a $\mathrm{C}_{1}-\mathrm{C}_{6}$ bond length intermediate between the two $\mathrm{C}_{2 \mathrm{~h}}$ minima on the CASSCF PES. This single $\mathrm{C}_{2 n}$ stationary point is the transition state for the Cope rearrangement; and the calculated enthalpy of activation is within about 1 $\mathrm{kcal} / \mathrm{mol}$ of experiment. There is little qualitative resemblance between the CASSCF results and those obtained from calculations which include dynamic electron correlation. ${ }^{43}$

Other Pericyclic Reactions. The Cope rearrangement is not the only pericyclic reaction for which one must go beyond the CASSCF level in order to correctly predict the reaction pathway. For example, in calculations on the Diels-Alder reaction at the

[^8]CASSCF/6-31G* level, Li and Houk found that a diradical intermediate, with just one $\mathrm{C}-\mathrm{C}$ bond formed between ethylene and butadiene, is $3.1 \mathrm{kcal} / \mathrm{mol}$ lower than the concerted transition state. ${ }^{18}$ In contrast, at the RQCISD(T) level, in which all the valence electrons are correlated, the concerted transition state is $4.3 \mathrm{kcal} / \mathrm{mol}$ lower in energy than the diradical intermediate and $10.2 \mathrm{kcal} / \mathrm{mol}$ below the transition state leading to this intermediate. Moreover, the energy of the concerted transition state, relative to the reactants, drops from $43.8 \mathrm{kcal} / \mathrm{mol}$ at the CASSCF level to $25.5 \mathrm{kcal} / \mathrm{mol}$ at the RQCISD(T) level. The latter is very dose to the activation energy measured for this reaction. ${ }^{18}$

For the ring opening of cyclobutene to butadiene, CASSCF and CASPT2N calculations with the 6-31G* basis set give, respectively, 36.6 and $36.3 \mathrm{kcal} / \mathrm{mol}$ for the energy difference between cyclobutene and the conrotatory transition state. ${ }^{17}$ These values are both close to the "experimental" value of $34.5 \mathrm{kcal} / \mathrm{mol} .{ }^{44}$ Therefore, it might appear that CASSCF is adequate for this reaction and that inclusion of dynamic electron correlation is unnecessary.
That this is not the case is shown by the calculated energy difference between the transition state and the product, transoid-1,3-butadiene. This difference is $55.4 \mathrm{kcal} / \mathrm{mol}$ at the CASSCF level and $46.9 \mathrm{kcal} / \mathrm{mol}$ at the CASPT2N level. The "experimental" energy difference is $45.4 \mathrm{kcal} / \mathrm{mol}$. ${ }^{44}$
CASSCF gives a satisfactory value for the energy difference between cyclobutene and the transition state for ring opening because (4/4)-CASSCF provides correlation for only one of the four strained $\sigma$ bonds in cyclobutene. This results in the overestimation of the energy of cyclobutene, relative to that of 1,3butadiene, by about $8 \mathrm{kcal} / \mathrm{mol}$. The CASSCF overestimation of the energy of cyclobutene balances the $10 \mathrm{kcal} / \mathrm{mol}$ overestimation of the energy of the transition state for conrotatory ring opening, caused by the absence of dynamic electron correlation in the CASSCF wave function.

## Dehydrobenzenes

In 1,2- and 1,3-dehydrobenzene ( $\mathbf{1}$ and $\mathbf{2}$ ) the additional bonding, not present in benzene, between the "dehydrocarbons" is rather weak. For example, TC-SCF/4-31G calculations by Noell and Newton found the energy of $\mathbf{1}$ and $\mathbf{2}$ to be, respectively, only 23.3 and $8.8 \mathrm{kcal} / \mathrm{mol}$ lower than that of 1,4 -dehydrobenzene (3). ${ }^{45}$ In 3 the through-space interaction between the dehydrocarbons is so weak that the dominant interaction between them is through the $\mathrm{C}_{2}-\mathrm{C}_{3}$ and $\mathrm{C}_{5}-\mathrm{C}_{6} \sigma$ bonds.

TCSCF calculations with basis sets much better than 4-31G give similar results. ${ }^{19}$ TheTCSCF results are in good agreement with the first set of experimental results on the relative heats of formation of these three isomers, published by Wenthold, Paulino, and Squires. ${ }^{20}$

The TCSCF wave functions correlate the two electrons in the weak $\sigma$ bond between each pair of dehydrocarbons in 1-3, but these wave functions provide no correlation between the two $\sigma$ electrons and the six $\pi$ electrons. Inclusion of the latter type of

[^9]correlation should be important, since the required $\sigma$ $\rightarrow \sigma^{*}, \pi \rightarrow \pi^{*}$ excitations are relatively low in energy. This type of correlation would be expected to stabilize ionic terms in the wave functions for the two electrons in the weak bonds between the dehydrocarbons in 1 and 2, thus increasing the occupancy of the bonding $\sigma \mathrm{MOs}$ and resulting in stronger bonds between these carbons.

Calculations that include dynamic correlation between the $\sigma$ and $\pi$ electrons have found this to be the case. ${ }^{19}$ The weight of the $\left|\ldots \sigma^{* 2}\right\rangle$ configuration, relative to that of $\left|\ldots \sigma^{2}\right\rangle$, is smaller in these wave functions than in those computed at the CASSCF level; and the heats of formation of $\mathbf{1}$ and $\mathbf{2}$ are caclulated to be, respec-
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tively, 28-31 and 13-14 kcal/mol lower than that of 3.

The disagreement between the results of these calculations and the first set of experimental results ${ }^{20}$ on the heats of formation of 1-3 led Wenthold and Squires to identify a problem in these experiments and to devise a very clever way to circumvent it. ${ }^{21}$ A second set of experiments gave heats of formation for 1-3 that are in excellent agreement with those predicted by the calculations that included dynamic electron correlation. ${ }^{19}$

## Conclusions

A great deal of conceptual simplicity is provided by descriptions of molecules and reactions which focus on just the few electrons that are "active". However, in order to obtain satisfactory results in ab initio calculations on excited states that have ionic character, radical ions, pericyclic reactions, and molecules that contain weak bonds, one cannot focus on just the active electrons. ${ }^{46}$ Instead, one must go beyond the CASSCF level and include dynamic correlation between the "active" and "inactive" electrons.

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