# Effects of Electron Correlation and Spin Projection on Rotational Barriers of Trithiocarbenium Ion $[C(SH)_3]^+$ and Radical Dication $[C(SH)_3]^{\cdot,2+}$

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Received 19 July 1996; accepted 14 October 1996

**ABSTRACT:** Theoretical level dependencies are discussed of relative isomer stabilities and rotational barriers of trithiomethyl cation  $[C(SH)_3]^+$  (a) and of radical dication  $[C(SH_3)]^{+,2+}$  (b). Spin polarization and dynamic electron correlation are very important for the radical dictation. Removal of an electron from one of the degenerate  $\pi$ -HOMOs of C<sub>3h</sub> symmetric [C(SH)<sub>3</sub>]<sup>+</sup> stabilizes the remaining  $\pi$  electron to such an extent that the unpaired electron is not in the HOMO of the dictation. The radial  $\pi$  MO's "diving below the Fermi level" facilitates strong spin polarization because of its energetic proximity to  $\sigma$  MOs. Projection of the first three higher spin states eliminates spin contaminations, and the terms E(PUHF(s + 3)) - E(UHF) and E(PMP4(s + 3)) - E(MP4) are discussed. The combination of annihilation of spin contamination and electron correlation is essential for the determination of relative energies and rotational barriers of the radical dication. The results obtained at this level match the results of high level QCISD(T) calculations in a near-quantitative fashion. Perturbation theory alone does not correct for spin contamination even if it is carried to full fourth order and includes triple excitations; the E(PMP4(s + 3))-E(MP4) values are all negative and can exceed 5 kcal/mol in magnitude. Previous studies showed that annihilation of spin contaminations is important in regions of potential energy surfaces where  $\sigma$  bonds are broken (homolytic dissociation), formed (radical addition), or both (H abstraction by radical). Our findings stress that the annihilation of spin contaminations can be just as

Dedicated to Prof. Andrew Streitwieser on the occasion of his 70th birthday.

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Contract grant sponsor: NATO

important for any process that greatly alters spin polarization and even if that process proceeds without breaking or forming of  $\sigma$  bonds. For comparison, density functional theory also was employed in the potential energy surface analyses. The results obtained with the B3LYP formalism were found to be less susceptible to spin contamination and resulted in rather good agreement with the best pertubation and configuration interaction results. © 1997 by John Wiley & Sons, Inc. *J Comput Chem* **18**: 1023–1035, 1997

**Keywords:** relative isomer stabilities; rotational barriers; thiomethyl cation; radical dication; electron correlation; spin projection

# Introduction

ulfur is effective at stabilizing polycations<sup>1</sup> due to its low ionization potential and electronegativity.<sup>2</sup> Thiocarbenium ions<sup>3</sup> were prepared and characterized<sup>4-6</sup> and some of these are stable in the solid state. X-ray crystal structures of several di-7 and trithiosubstituted<sup>8</sup> carbenium ions were obtained. In a recent study that combined experimental and ab initio computational investigations of the systems  $[CH_n(XR)_{3-n}]^+[X = O, S]$ Se, Te;  $R(exp.) = 2, 4, 6-i Pr_3C_6H_2$ , R(calcd.) = H; n = 0-2], we found that the mode of stabilization by sulfur and the heavier homologs differs greatly from the mechanism of stabilization of carbenium ions by oxygen.9 These studies indicated an enhanced charge separation in [C(OH)<sub>3</sub>]<sup>+</sup> instead of the commonly expected charge delocalization. For  $[C(OH)_3]^+$ , natural population analysis (NPA) suggested a positive charge on carbon that is larger than unity (+1.3). The homologous ions  $[C(XH)_3]^+$ (X = S, Se, Te) were found to differ fundamentally in that the carbon center is negatively charged as the result of effective  $X \rightarrow C$  donation in the  $\sigma$ and  $\pi$  systems. In a subsequent study, we explored the possibility of further oxidation for the sulfur system via ab initio potential energy surface analysis of  $[C(SH)_3]^+$  and of the radical dication  $[C(SH)_3]^{,2+,10}$  The electronic features suggested by the NPA analysis for the monocation were fully corroborated by topological analysis of the electron density distributions in Cartesian space. Moreover, it was found that this unexpected electronic motif persists even in the dication. Oxidation of the monocation primarily removes sulfur  $\pi$ -electron density and removes only some electron density from the central C atom. The  $\alpha$ -excess spin density is entirely concentrated on S atoms and the carbon carries some  $\beta$ -spin density because of spin polarization. Minima and transition state structures for isomerization were reported for  $[C(SH)_3]^+$  and  $[C(SH)_3]^{\cdot,2+}$  and (P)MP4(full, sdtq)/ 6–31G\*// HF/ 6–31G\* +  $\Delta$ VZPE(RHF/6–31G\*) level activation barriers to rotation were discussed as probes for the importance of conjugation.<sup>11</sup>

With the present study we expand on one aspect of our studies of the cations  $[C(SH)_3]^{n+}$  that is of more general theoretical interest, and this aspect concerns the energetic effects of annihilation of unwanted spin contaminations in the wave functions of the radical [C(SH)<sub>3</sub>]<sup>·,2+</sup> at the Hartree-Fock (HF) and Møller-Plesset (MP) levels. In 1986 Schlegel combined unrestricted fourth-order perturbation theory with spin projection (PMP4) to annihilate the largest spin contaminant,<sup>12</sup> and the methodology for full annihilation was developed shortly thereafter.<sup>13</sup> Homolytic dissociations were studied and approximate and complete annihilation gave PMP4 potential energy curves that agreed well with very accurate configuration interaction (CI) calculations. The potential energy surface characteristics can be affected significantly in regions where spin contamination becomes important and especially when electron correlation corrections are treated by MP perturbation theory. In one study of the addition of an H atom to ethyne, ethene, and formaldehyde, the effects of spin contamination on both geometries and activation barriers were studied at the PMP2 level. The PMP2 data provided a great improvement over the MP2 level and closely approximated the PMP4 results.<sup>14</sup> Studies of radical additions to simple unsaturated compounds<sup>15–17</sup> showed that good agreement with an experiment can be obtained when corrections for spin projection are combined with electron correlation corrections while the barrier heights generally were too high when spin contamination was ignored. Subsequent studies (Table I) of the addition of ethene radical cation to ethene,<sup>18</sup> of the additions of tert-butyl radical and benzyl radical to ethene,<sup>19</sup> and of the addition of simple  $\sigma$  radicals to mono- and difluoroethene<sup>20</sup> corroborated

Method	Systems Studied	Authors, Year
PMP4, PA of SC	Diss. of LiH of HF	Schlegel, 1986 <sup>12</sup>
PMP4, CA of SC	Diss. of LiH, HF, and $H_2O$	Schlegel, 1988 <sup>13</sup>
PMP2 structures	Add. of H <sup><math>\cdot</math></sup> to C <sub>2</sub> H <sub>2</sub> , C <sub>2</sub> H <sub>4</sub> & H <sub>2</sub> CO	McDouall & Schlegel, 1989 <sup>14</sup>
PMP4, PA of SC	Add. of HO <sup><math>\cdot</math></sup> to $\tilde{C}_{2}H_{2}$ & $\tilde{C}_{2}H_{4}$	Sosa & Schlegel, 1987 <sup>15</sup>
PMP4, PA of SC	Add. of H <sup><math>\cdot</math></sup> to C <sub>2</sub> H <sub>4</sub> & H <sub>2</sub> CO <sup><math>+</math></sup>	Sosa & Schlegel, 1986 <sup>16</sup>
PMP4, PA of SC	Add. of Me <sup>+</sup> to $\tilde{C}_2H_4 \& H_2CO$	Gonzalez et al., 1989 <sup>17</sup>
PMP4, CA of SC	H abstr. from $CH_{4}$ by H, $HO$ ; or $O_{3}$	Gonzalez et al., 1990 <sup>22</sup>
PMP4, PA of SC	<sup>2</sup> N and <sup>4</sup> N reactions with Me	Gonzalez & Schlegel, 1992 <sup>26</sup>
PMP4, PA of SC	H abstr. from HXCO (X = H, F, Cl) by HO <sup><math>\cdot</math></sup>	Francisco, 1992 <sup>23</sup>
PMP4, CA of SC	Add. of $C_4 H_4^{+}$ to $C_2 H_4$	Alvarez-Idaboy et al., 1993 <sup>18</sup>
PMP4, PA of SC	H abstr. from NH <sub>3</sub> by HO	Corchado et al., 1993 <sup>24</sup>
PMP4, CA of SC	Add. of tert- $C_{4}H_{0}^{\dagger}$ and Bz $^{\circ}$ to $C_{2}H_{4}$	Arnaud et al., 1994 <sup>19</sup>
PMP4, CA of SC	Add. of $\sigma$ radicals to fluoroethene	Donovan & Famini, 1994 <sup>20</sup>
PMP4, PA of SC	Add. of bicyclo [2,1,0]pentyl radical	Bach et al., 1994 <sup>27</sup>
PMP2, PA of SC	Add of R to alkenes	Wong & Radom, 1995 <sup>21</sup>
PMP4, CA of SC	H abstr. from chloroethane by HO	Sekusak et al., 1995 <sup>25</sup>

#### TABLE I. \_\_\_\_\_ Development and Applications of Spin Projected Møller–Plesset Perturbation Theory.

<sup>a</sup> Spin projected fourth-order Møller-Plesset theory (PMP4) with partial (PA) or complete annihilation (CA) of spin contaminants (SC).

that the energetics of the radical addition reactions can be very sensitive to electron correlation and spin contamination. Recent theoretical model dependencies of some prototypical radical addition reactions further emphasize this point.<sup>21</sup> While the thermochemistry of the reactions shows moderate theoretical level dependency, significant differences of up to 10 kcal/mol were found between activation barriers computed with and without annihilation of spin contaminations. Importantly, the work of Alvarez-Idaboy et al.18 shows that elimination of spin contaminants can increase the activation barriers while in most cases a decrease of activation barriers is observed. The study by Donovan and Famini<sup>20</sup> is important because it shows that the predicted regiochemistry can depend on whether the annihilation of spin contaminants was or was not combined with a perturbational electron correlation treatment. Similar methodological effects were reported for H-abstraction reactions from methane,<sup>22</sup> from HXCO (X = H, F, Cl)<sup>23</sup> from ammonia,<sup>24</sup> and from chloroethane.<sup>25</sup> In the study by Covchado et al.,<sup>24</sup> for example, it was pointed out that the spin projection had larger effects on the kinetic barrier than on the reaction thermochemistry and the PMP4 calculations provided activation barriers for H transfer that agree well with the experiments. These same conclusions are true for other H-abstraction reactions and for the insertion reaction as well.<sup>26</sup> Yet, there are again some notable exceptions. Recently, Bach et al. studied the properties of the radical obtained by H abstraction from bicyclo-[2.1.0]pentane and its reactions (H abstraction, HO' addition) at the MP4 level without and with annihilation of spin contamination. The activation barriers determined at the MP4 and PMP4 levels differed only very slightly.<sup>27</sup>

All of these studies of annihilation of spin contaminations (vide supra) can be summarized by stating that spin projection becomes important in regions of the potential energy surfaces where  $\sigma$ bonds are broken (homolytic dissociation), formed (radical addition), or both (H abstraction). Because the annihilation of spin contaminants reduces spin polarization,<sup>28</sup> one would expect the largest effects in the transition regions because these regions are characterized by the most spin decoupling and therefore are prone to realize strong spin polarization. We have been interested in spin density distributions and recently reported a study on the rotational automerization of allyl radical.<sup>28b</sup> It was found that spin projection was necessary to predict the correct transition state structure. Only when perturbational corrections to electron correlation were combined with annihilation of spin contamination did the predicted activation barrier for rotational automerization agree with high level CI data and with the experiment. Our study of allyl radical shows that the effects of spin contamination can be significant even in processes in which no  $\sigma$  bonds are broken and where the electronic reorganization merely involves the breaking of  $\pi$ dative bonds and spin localization. This conclusion is corroborated by other studies of the allyl radical<sup>29</sup> as well as the studies of the benzyl radical.<sup>30</sup> Here, we present a comparative analysis of relative isomer stabilities and of rotational barriers of  $[C(SH)_3]^+$  and  $[C(SH)_3]^{\cdot,2+}$ . This analysis has several attractive features to add to this discussion. The theoretical level affects the monocation characteristics only slightly and this system provides a good point of reference. A most pertinent advantage relates to the fact that dynamic electron correlation becomes particularly important for the radical dication. It is known that dynamic electron correlation between  $\pi$  and  $\sigma$  electrons is critical in molecules with charged  $\pi$  systems,<sup>31</sup> and this is even more true for the dication for the following reason. Removal of one electron from one of the degenerate  $\pi$  highest occupied molecular orbitals (HOMOs) of  $[C(SH)_3]^+$  stabilizes the remaining  $\pi$ electron greatly and the MO of the unpaired  $\pi$ radical is not the HOMO in the dication. The unpaired  $\pi$  MOs "diving below the Fermi level" facilitates strong spin polarization because of its energetic proximity to  $\sigma$  MOs. It is for this scenario that energetic effects of spin contamination and its annihilation should be particularly pronounced, even though no  $\sigma$  bonds are broken or formed. It will be shown that the combination of annihilation of spin contamination and electron correlation is essential for the determination of relative isomer stabilities and for activation barriers. At this level, the results of the perturbation calculations match high level QCIS(T) data in near-quantitative fashion. Quadratic CI theory is much less sensitive to spin contamination and thus can serve as the standard for the assessment of the accuracy of the perturbation results. Much less is known about the reliability of density functional theory (DFT) in such cases, and we present results of semiempirical solutions for comparison.

# **Theory and Computations**

Restricted (RHF) and unrestricted (UHF) Hartree–Fock or HF theory<sup>32</sup> were employed for the closed- and open-shell systems, respectively. While the wave functions obtained with the UHF formalism are eigenfunctions of the Hamiltonian and the  $S_z$  operators,<sup>33</sup> they are not eigenfunctions of the  $S^2$  operator. As a result, the wave functions

of the doublet systems are spin contaminated to some extent by admixtures of quartet, sextet, and higher spin states.<sup>34,35</sup> The eigenvalues of the  $S^2$ operator are given as a measure of the spin contamination. Because the projection operator commutes with the charge density operator,<sup>36</sup> the electron density remains unaffected by the projection and effects of spin contamination on the optimized structures are expected to be small.<sup>28</sup> Complete gradient optimizations of geometries and vibrational analysis were carried out at the HF/6-31G\* level as described previously.<sup>10</sup> The zero-point energies calculated at the HF level were scaled in the usual fashion<sup>37</sup> (factor 0.9) when applied to relative energies. Electron correlation effects on relative stabilities were estimated using full fourthorder MP perturbation theory [MP4(full, sdtq)/6-31G\*//HF/6-31G\*]. Borden and Davidson stressed the importance of triple excitations to correctly compute molecules with charged  $\pi$  systems.<sup>31</sup> The work by Knowles and Handy<sup>38</sup> points out that spin contamination adversely affects the convergence of the perturbation series and that a significant improvement can be obtained after spin projection. To probe these energetic consequences of spin contamination, we also determined the HF and MP4 energies after annihilation. QCI theory<sup>39</sup> with all single and double excitations considered and including a perturbation correction for triple excitations, QCISD(T), is used as the high level standard in the assessment of the performances of all other methods. DFT<sup>40</sup> has emerged in recent years as one of the more promising approaches within the arsenal of semiempirical methods and we have included the results of B3LYP<sup>41</sup> calculations for comparison. In Table II we report the results of the potential energy surface exploration at the HF level and higher level energies determined with the  $HF/6-31G^*$  structures. The results of computations that involved the annihilation of spin contaminants are listed in Table III. The energies PUHF(s + 1) and PUHF(s + 3) were obtained after removal of the next higher spin contamination (quartet) and after removal of the spin contaminations associated with the first three higher spin states (s + 1 to s + 3) Projection of just the next higher spin state results in  $S^2$  eigenvalues of  $0.750 \pm 0.001$  and only for **1b** and **2b** the  $S^2$ values of  $0.750 \pm 0.060$  remain slightly higher. Annihilation of the first higher spin state removes the major part of the spin contamination successfully as indicated by the  $S^2$  eigenvalues after projection of the quartet state.<sup>42,43</sup> Projection of the first three

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	Sym.	HF/6-31G*			MP4(sdta)	B3LYP	QCISD(T)
Molecule		Energy	VZPE	NIMAG	Energy	Energy	Energy
C(SH) <sub>3</sub> +, <b>1a</b>	C <sub>3h</sub>	- 1231.854177	24.27	0	- 1232.450046	- 1234.174283	- 1232.450180
C(SH) <sub>3</sub> <sup>2+</sup> , <b>1b</b>	C <sub>3h</sub>	- 1231.319016	23.16	0	- 1231.888288	- 1233.608296	- 1231.898812
C(SH) <sub>3</sub> <sup>2+</sup> , 1b'	Cs	- 1231.319912	23.19	0	- 1231.890024	- 1233.609855	- 1231.900278
C(SH) <sub>3</sub> <sup>+</sup> , <b>2a</b>	Cs	- 1231.850775	24.16	0	- 1232.446974	- 1234.171133	- 1232.447104
C(SH) <sub>3</sub> <sup>2+</sup> , <b>2b</b>	Cs	- 1231.321188	23.28	0	- 1231.891893	- 1233.610912	- 1231.901983
C(SH) <sub>3</sub> <sup>2+</sup> , <b>3b-c1</b>	C <sub>1</sub>	- 1231.240092	23.86	0	- 1231.827837	- 1233.544650	- 1231.830816
C(SH) <sub>3</sub> <sup>2+</sup> , <b>3b-c2</b>	C <sub>1</sub>	- 1231.243485	24.18	0	- 1231.830563	- 1233.546176	- 1231.833077
C(SH) <sub>3</sub> <sup>2+</sup> , <b>3b-t1</b>	C <sub>1</sub>	- 1231.245134	24.07	0	- 1231.831865	- 1233.547899	- 1231.834497
C(SH) <sub>3</sub> <sup>2+</sup> , <b>3b-t2</b>	C <sub>1</sub>	- 1231.245215	24.10	0	- 1231.831733	- 1233.548147	- 1231.834548
C(SH) <sub>3</sub> <sup>+</sup> , <b>4a</b>	C <sub>1</sub>	- 1231.837539	23.89	1	- 1232.434893	- 1234.157807	- 1232.435292
C(SH) <sub>3</sub> <sup>2+</sup> , <b>4b</b>	C <sub>1</sub>	- 1231.310806	23.01	1	- 1231.892639	- 1233.601374	- 1231.894806
C(SH) <sub>3</sub> <sup>+</sup> , <b>5a-u</b>	Cs	- 1231.835657	23.96	1	- 1232.433210	- 1234.156099	- 1232.433562
C(SH) <sub>3</sub> <sup>+</sup> , <b>5a-w</b>	Cs	- 1231.836563	23.84	1	- 1232.434070	- 1234.156970	- 1232.434513
C(SH) <sub>3</sub> <sup>2+</sup> , <b>5b-u</b>	Cs	- 1231.313212	23.10	1	- 1231.894728	- 1233.600774	- 1231.895788
C(SH) <sub>3</sub> <sup>2+</sup> , <b>5b-w</b>	Cs	- 1231.305256	22.95	1	- 1231.889498	-1233.598623	- 1230.891268

#### TABLE II. \_\_\_\_\_\_ Total and Vibrational Zero-Point Energies.

Energies in atomic units.  $RHF/6-31G^*$  for closed shell systems and  $UHF/6-31G^*$  for open shell systems. MP4(sdtq) and QCISD(T) calculations included all electrons in the active space. The MP4(sdtq), QCISD(T), and B3LYP calculations employed the 6-31G\* basis set and were based on the HF/6-31G\* structures. Vibrational zero-point energies (VZPE) in kcal/mol as calculated. NIMAG, number of imaginary frequencies.  $S^2$  eigenvalues of UHF wave functions: **1b**, 1.1431; **1b**, 1.1410; **2b**, 1.1120; **3b-c1**, 0.7766; **3b-c2**, 1.1761; **3b-t1**, 0.7734; **3b-t2**, 0.7770; **4b**, 0.7883; **5b-u**, 0.7630; **5b-w**, 0.7631.

# TABLE III. Results of Calculations Involving Annihilation of Spin Contamination.

•		•			
PUHF(s + 1) Energy	$\Delta E_1$	PUHF(s + 3) Energy	$\Delta E_2$	PMP4(s + 3) Energy	$\Delta E_3$
- 1231.336728	-11.11	- 1231.335311	0.89	- 1231.900634	-7.75
- 1231.337373	- 10.96	- 1231.335983	0.87	- 1231.902175	-7.62
- 1231.338129	- 10.63	- 1231.336884	0.78	- 1231.903563	-7.32
<b>1</b> – 1231.245042	-3.11	- 1231.245000	0.03	- 1231.829719	-1.18
<b>2</b> – 1231.248346	-3.05	- 1231.248316	0.02	- 1231.832228	-1.04
<b>1</b> – 1231.249867	-2.97	- 1231.249837	0.02	- 1231.833493	-1.02
<b>2</b> – 1231.250162	-3.10	- 1231.250123	0.02	- 1231.833631	-1.19
- 1231.315029	-2.65	- 1231.317971	0.04	- 1231.895114	- 1.55
– 1231.316630	-2.14	- 1231.316623	0.00	- 1231.895847	-0.70
v -1231.308672	-2.14	- 1231.308665	0.00	- 1231.890627	-0.71
	PUHF(s + 1) Energy - 1231.336728 - 1231.337373 - 1231.338129 1 - 1231.245042 2 - 1231.245042 2 - 1231.248346 1 - 1231.249867 2 - 1231.250162 - 1231.315029 1 - 1231.316630 4 - 1231.308672	$\begin{array}{c} {\sf PUHF(s+1)} \\ {\sf Energy} & \Delta E_1 \\ \\ -1231.336728 & -11.11 \\ -1231.337373 & -10.96 \\ -1231.338129 & -10.63 \\ 1 & -1231.245042 & -3.11 \\ 2 & -1231.245042 & -3.11 \\ 2 & -1231.248346 & -3.05 \\ 1 & -1231.249867 & -2.97 \\ 2 & -1231.250162 & -3.10 \\ -1231.315029 & -2.65 \\ 1 & -1231.316630 & -2.14 \\ 4 & -1231.308672 & -2.14 \\ \end{array}$	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $

Energies in atomic units.  $\Delta E$  values in kcal/mol.

#### TABLE IV. \_\_\_\_\_ Relative Energies.

Parameter	HF No Proj.	∆VZPE Scaled	PUHF Proj. s + 1	PUHF Proj. s + 3	MP4 No. Proj.	PMP4 Proj. s + 3	DFT B3LYP	QCISD(T)
I-Pref (1b' over 1b)	0.56	-0.03	0.40	0.42	1.09	0.97	0.98	0.92
I-Pref ( <b>1a</b> over <b>2a</b> )	2.13	-0.10			1.93		1.98	1.93
I-Pref (1b over 2b)	- 1.36	0.11	-0.88	-0.99	-2.26	- 1.84	-1.64	- 1.99
I-Pref (3b-t2 over 3b-t1)	0.05	-0.03	0.19	0.18	-0.08	0.09	0.16	0.03
I-Pref (3b-t2 over 3b-c2)	1.09	0.07	1.14	1.13	0.73	0.88	1.24	0.92
I-Pref (3b-c2 over 3b-c1)	2.13	-0.29	2.07	2.08	1.71	1.57	0.96	1.42
I-Pref (1b over 3b-t2)	46.31	0.85	54.32	53.46	35.49	42.05	37.74	40.33
$E_A(1a \Rightarrow 4a)$	10.44	-0.34			9.51		10.34	9.34
$E_A(1b \Rightarrow 4b)$	5.15	-0.13	13.62	12.76	-2.73	3.46	4.34	2.51
$E_A(2a \Rightarrow 4a)$	8.31	-0.24			7.58		8.36	7.41
$E_A(\mathbf{2b} \Rightarrow \mathbf{4b})$	6.51	-0.24	14.50	13.75	-0.47	5.30	5.99	4.50
$E_A$ (2a $\Rightarrow$ 5a-u)	9.49	-0.18			8.64		9.43	8.50
$E_A(2a \Rightarrow 5a-w)$	8.92	-0.29			8.10		8.89	7.90
$E_A(2b \Rightarrow 5b-u)$	5.01	-0.16	13.49	12.71	-1.78	4.84	6.36	3.89
$E_A(\mathbf{2b} \Rightarrow \mathbf{5b-w})$	10.00	-0.30	18.48	17.71	1.50	8.12	7.71	6.72

Isomer preference energies (I-Pref), activation energies ( $E_A$ ), ionization energies (IE), and hydride affinities (HA) in kcal/mol.  $\Delta$ VZPE terms calculated at the HF level are scaled (factor 0.9), and they are to be added to the relative energies.



**FIGURE 1.** Structures of the minima  $C_{3h}$ -1a and  $C_s$ -2a of  $[C(SH)_3]^+$ . Structures 1b  $(C_{3h})$  and 1b'  $(C_s)$  are minima of dication  $[C(SH)_3]^{\cdot,2+}$  resulting by oxidation of  $C_{3h}$ -1a. Dication  $C_s$ -2b is the oxidation product of 2a. Values given in italics for 1a are the average X-ray data for  $[C(SR)_3]^+$  with R = 2,4,6-tri-(isopropyl)phenyl.

unwanted spin states results effectively in complete annihilation. Pertinent relative energies obtained without and with annihilation at the HF and MP4 levels as well as at the DFT and QCI levels are summarized in Table IV. Calculations were performed with the program Gaussian 94<sup>44</sup> and earlier versions on a network of IBM/RS-6000, Silicon Graphics Indigo, and PowerChallenge L computers.

#### **Results and Discussion**

#### EQUILIBRIUM STRUCTURES AND ISOMERIZATIONS OF $[C(SH)_3]^+$ AND $[C(SH)_3]^{\cdot,2+}$

An analysis of the potential energy surfaces of  $[C(SH)_3]^+$  and  $[C(SH)_3]^{\cdot,2+}$  was given previously<sup>10</sup> and we briefly review the relevant parts. Isomeric structures are distinguished by numbers and the labels **a** and **b** are used to denote the cation and the dication, respectively. Unless otherwise noted, relative energies in the following refer to the highest level of perturbation theory, PMP4(full, sdtq)/6–31G\*//HF/6–31G\* +  $\Delta$ VZPE (HF/6–31G\*).

The  $C_{3h}$  structures **1a** and **1b** are minima (Fig. 1). The X-ray structure of  $[C(SR)_3]^+$  with R =2,4,6-tri(isopropyl)phenyl realizes de facto C<sub>3h</sub> symmetry of the central unit and the C-S bond lengths agree well with the calculated structure 1a. The wave function for 1b is asymmetric and allows for reduction in molecular symmetry. Optimization within C<sub>s</sub> symmetry leads to the Jahn-Teller distorted structure 1b', which is preferred over 1b by less than 1 kcal/mol. 1b' is modestly distorted in that two C-S bonds are elongated while one is shortened. The distorted structure **1b**" in which one C—S bond is elongated while the two other are shortened corresponds to the transition state structure for pseudorotation.<sup>45</sup> The rather similar energies of 1b and 1b' indicate that the pseudorotation of 1b' is essentially unhindred and there is no need to optimize 1b". Rotation about one of the C-S bonds in 1 leads to structures 2 and the C<sub>s</sub> structures 2a and 2b are minima. While 1a is preferred over 2a, the dication 1b is less stable than 2b. Several distonic stereoisomers with S-S bonds were located (Fig. 2). In 3b-c1 and 3b - c2, the endocyclic S—H bonds are cis with regard to each other, while the endocyclic S—H bonds are trans to each other in 3b-t1 and 3b-t2. Isomers 3b-t2 and 3b-t1 are nearly isoenergetic and the cis structures are



**FIGURE 2.** Stereoisomers **3b** are local minima of the radical dication  $[C(SH)_3]^{\cdot,2+}$ .

more stable than the trans structures. All of the isomers of **3b** are more than 35 kcal/mol less stable than  $C_{3h}$ -**1b**. The isomerization between **2** and **1** involves rotation about the c bond of **2** indicated in Figure 1 and involves the chiral transition state structures **4a** and **4b**. Rotation about the a or b bonds in **2a** or **2b** results in automerization via the  $C_s$ -symmetric transition state structures **5a** and **5b** (Fig. 3). The structures **5** come as isomers **5-w** and **5-u**, depending on whether rotation occurs about the a or the b bond, respectively. The labels w and u describe the shape of the in-plane H—S—C—S—H fragment.

#### **MO DIAGRAMS OF RADICAL DICATIONS**

The valence-MO diagrams of **1a** and of the oxidized species **1b** and **1b**' are shown in Figure 4. The RHF MO levels are shown for **1a** and for the radicals set of  $\alpha$ - and  $\beta$ -spin orbitals are shown. With the exception of  $\pi_{28}$  of the dication, all of the MOs are occupied but, for the sake of clarity, only the  $\pi$  electrons are indicated by  $\uparrow$  ( $\alpha$  spin) and  $\downarrow$  ( $\beta$  spin).



**FIGURE 3.** The transition state structures for isomerization  $1 \Rightarrow 4^{\ddagger} \Rightarrow 2$  and automerizations  $2 \Rightarrow 5 \cdot u^{\ddagger} \Rightarrow 2$  and  $2 \Rightarrow 5 \cdot w^{\ddagger} \Rightarrow 2$  are shown for the monocation (top) and the dication (bottom).

Chemists are inclined to assume that the oxidation of a closed-shell system leads to an open-shell system in which the unpaired electron resides in the HOMO. This assumption, of course, reflects an education in MO theory that for several decades used to focus on Hückel theory. Removal of one electron from the degenerate  $\pi$  HOMO ( $\pi_{27/28}$ ) results in a stabilization of the remaining unpaired  $\pi$  electron ( $\alpha$ - $\pi_{26}$ ) significantly below the energy of the remaining quasidegenerate  $\pi$  HOMO ( $\alpha$ - $\pi_{28}$ and  $\beta$ - $\pi_{27}$ ) of the radical dication. The energies of the electrons of the all-bonding  $\pi_{23}$  MO in **1a** become greatly different in the dication  $(\alpha - \pi_{23})$ and  $\beta$ - $\pi_{24}$ ). While the lowering of the MO of the unpaired electron below the energy of the remaining doubly occupied MO of the formerly degenerate HOMO set, although hardly ever recognized, can be rationalized as a consequence of a reduction of electron-electron repulsion, it is most intriguing indeed that sets of spin paired  $\pi$  and  $\sigma$  electrons are both higher in energy compared to the MO associated with the unpaired  $\pi$  radical. Dynamic electron correlation becomes so important in this charged  $\pi$  radical because of this readily identifiable feature of the MOs.

#### ENERGETIC EFFECTS OF ANNIHILATION OF SPIN CONTAMINATIONS AND OF ELECTRON CORRELATION

The energetic effects of annihilation of spin contaminations and of electron correlation are conveniently discussed with the parameters  $\Delta E$  included in Table III. The energy differences  $\Delta E_1 =$ E(PUHF(s + 1)) - E(UHF) and  $\Delta E_2 = E(PUHF(s + 1)) - E(UHF)$ + 3)) – E(PUHF(s + 1)) allow for an appreciation of partial and complete annihilation of spin contamination at the HF level. Effects of spin annihilation at the electron correlated level were assessed via  $\Delta E_3 = E(PMP4(s + 3)) - E(MP4)$  and only complete annihilation was considered in this case. A positive  $\Delta E$  means that decontamination increases the energy. As can be seen from Table III, the energy differences  $\Delta E_1$  are substantial and range from +5 to -10 kcal/mol. The  $\Delta E_2$  values are all positive and are generally small. Because



FIGURE 4. The MO diagrams for 1a, 1b, and 1b' are drawn to scale.

the contaminations are completely eliminated after projection of the first three higher spin states, we report the PMP4(s + 3) energies together with the  $\Delta E_3$  values. The  $\Delta E_3$  values all are negative; that is, the PMP4 energies are lower than the MP4 energies throughout; and the magnitudes of  $\Delta E_3$ are large for most of the open structures while being modest for the ring structures.

An example for significant deviation between MP4 and PMP4 data relates to the relative isomer preference between **1b** and the distonic structures **3b**. As can be seen from the  $\Delta E_i$  values in Table III, the effects of spin projection are similar and small for all isomers of **3b**. Not so for **1b**. The annihilation of spin contaminations increases the PUHF value of I-Pref (**1b** over **3b**–**t2**) by nearly 10 kcal/mol. Electron correlation affects this value in the opposite direction and more, such that the PMP4 value becomes 42.1 kcal/mol. The MP4 value is about 6.5 kcal/mol lower still because the energy increase associated with proper spin annihilation is not accounted for.

The energetic consequences of annihilation of spin contaminations and/or of electron correlation

on relative stabilities of isomers 1 and 2 and on rotational barriers are demonstrated in a compelling fashion with the graphical representations of the potential energy surfaces of  $[C(SH)_3]^+$  and  $[C(SH)_3]^{\cdot,2+}$  shown in Figure 5. In the top left in Figure 5, the potential energy diagram is represented as computed at the HF/6-31G\* +  $\Delta VZPE$ level and energetic consequences of either annihilation of spin contamination or of electron correlation are shown in the lower left and the upper right, respectively. On the bottom right are shown the energetics obtained after annihilation and electron correlation at the PMP4/6-31G\*//HF/  $6-31G^* + \Delta VZPE$  level. In Figure 6 the results of the DFT and QCISD(T) calculations are presented in exactly the same manner as with Figure 5. It is immediately obvious that the best perturbation theoretical data agree in a near-quantitative fashion with the high-level QCISD(T) results. The results of the semiempirical density functional computations also agree quite well with these highly correlated *ab initio* data. While the differences between the correlated PMP4 level and the HF level are relatively small, very significant quantitative and even qualitative differences occur; however, if either spin contaminations are annihilated or electron correlation effects are included in the absence of accounting for the other effect. After a discussion of some characteristics of the profiles shown in Figure 5, a few instructive cases will be cited to highlight the methodological differences.

At the HF level, activation barriers of 10.1 and 5.0 kcal/mol are found for the processes  $1a \Rightarrow 4a^{\ddagger}$ and  $\mathbf{1b} \Rightarrow \mathbf{4b}^{\ddagger}$ , respectively. The rotational barrier are slightly reduced after annihilation of spin contamination and accounting for electron correlation effects. At the PMP4 level, the activation barriers for the processes  $1a \Rightarrow 4a^{\ddagger}$  and  $1b \Rightarrow 4b^{\ddagger}$  are 9.2 and 3.3 kcal/mol, respectively. This computed rotational barrier for 1a is comparable to the activation barriers of 8-14 kcal/mol measured for some disubstituted systems [RC(SR)2]<sup>+,7</sup> For the automerizations  $2a \Rightarrow 5a \cdot u^{\ddagger} \Rightarrow 2a$  and  $2a \Rightarrow 5a \cdot w^{\ddagger}$  $\Rightarrow$  2a, the activation barriers are 8.5 and 7.8 kcal/mol, respectively, at the highest level. For the automerizations of 2b via  $5b-u^{\ddagger}$  or  $5b-w^{\ddagger}$ , the activation barriers are 4.7 and 7.8 Kcal/mol, respectively. One would have expected that the rotational processes in the dication are less hindered than in the monocation and this is true for the isomerizations via 4 and for most automerizations but not for the automerization via **5b-w**<sup>‡</sup>. The automerization of 1 involves a sequence of isomerizations to 2 via  $4^{\ddagger}$  and automerizations of 2 via  $5^{\ddagger}$ ,





FIGURE 6. Density functional theory, B3LYP, and high level quadratic CI theory, QCISD(T), both agree very well with the scenario predicted at the PMP4 level. As in Figure 5, the potential energy diagram computed at the B3LYP/6-31G\*//HF/6-31G\* + ΔVZPE(HF/6-31G\*) level is represented on the left and the QCISD(T) / 6 −31G\* / / HF / 6 −31G\* + ∆VZPE(HF / 6 −31G\*) data are depicted on the right

and the former isomerizations are rate limiting for the monocation while the latter is rate limiting for the dication.

To illustrate the energetic effects of spin projection and electron correlation, a consideration of the automerization of 2b via 5b-u is particularly instructive. For the processes  $2a \Rightarrow 5a-u^{\ddagger} \Rightarrow 2a$  and  $2b \Rightarrow 5b-u^{\ddagger} \Rightarrow 2b$ , the activation barriers are 9.3 and 4.9 kcal/mol, respectively, at the HF level and the respective PMP4 barriers are similar (vide *supra*). Although **5b-u** is a transition state structure on the UHF potential energy surface, the MP4 energies calculated with the spin-contaminated UHF reference wave function would suggest that **5b-u** is more stable than **2b**. The MP2 level rotational profile of benzyl radical presents a similar case of such an artificial negative rotational barrier scenario.<sup>30</sup> With the data in Table IV we can trace the origin of this artifact. 2b is spin contaminated substantially, and more so than 5b-u, and the removal of these spin contaminants increases  $E_A(2b \Rightarrow 5b-u)$  from 5.0 to 12.7 kcal/mol primarily because 2b is stabilized by projection more than **5b-u** ( $\Delta E_1$  and  $\Delta E_2$  in Table III). On the other hand, electron correlation effects stabilize 5b-u more than 2b, counteracting the energetic effects of spin projection, and a PMP4 value results for  $E_4(2\mathbf{b} \Rightarrow 5\mathbf{b} \cdot \mathbf{u})$  that is close to the UHF value. The MP4 energies calculated without spin projection also indicate a reduction of  $E_A(2\mathbf{b} \Rightarrow 5\mathbf{b} \cdot \mathbf{u})$ ; but because spin annihilation, which increases  $E_A$ , is not properly accounted for, the resulting value of  $E_A(2\mathbf{b} \Rightarrow 5\mathbf{b} \cdot \mathbf{u}) = -1.78$  is much too low and would even suggest a change in the characteristics of the potential energy surface. This example well illustrates that MP perturbation theory, even when carried to full fourth-order, does not eliminate the spin contamination of the UHF reference and that PMP4 theory is required in such a case. Similar discussions apply to the activation barriers for the automerization  $2b \Rightarrow 5b$ -w and the isomerization  $1b \Rightarrow 4b \Rightarrow 2b.$ 

# Conclusion

Spin polarization and dynamic electron correlation are both important for the radical dication  $[C(SH)_3]^{\cdot,2+}$ . MO diagram analysis reveals the readily identifiable feature responsible for these effects. Removal of an electron from one of the degenerate  $\pi$  HOMOs of  $C_{3h}$  symmetric  $[C(SH)_3]^+$ results in a stabilization of the remaining  $\pi$  electron to such an extent that the unpaired electron is not in the HOMO of the dication. The "diving below the Fermi level'' of the unpaired  $\pi$  electron facilitates strong spin polarization because of the energetic proximity with  $\sigma$  MOs. The combination of annihilation of spin contamination and electron correlation is essential for the determination of relative energies and rotational barriers and affords an accuracy that matches the QCISD(T) results in a near-quantitative fashion. Importantly, pertubation theory alone does not correct for spin contamination even if it is carried to full fourthorder and includes triple excitations; the  $\Delta E_2$  values are all negative and can exceed 5 kcal/mol in magnitude. Previous studies showed that annihilation of spin contaminations is important in regions of potential energy surfaces where  $\sigma$  bonds are broken (homolytic dissociation), formed (radical addition), or both (H abstraction by radical). The results presented here stress that the annihilation of spin contaminations can be just as important for any process that greatly alters spin polarization and even if that process proceeds without breaking or forming of  $\sigma$  bonds.

# Acknowledgments

This research was supported by a NATO Collaborative Research Grant. We thank the MU Campus Computing Center for generous allocations of computer time and Hossein Tahani for his assistance.

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