## The Energetics of Valence Isomerization in the Norbornadiene–Quadricyclane System

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Potential applications of the photochromic valence isomerization of the norbornadiene(N)-quadricyclane(Q) system toward photochemical energy storage<sup>1</sup> and optical memory systems<sup>2</sup> have prompted numerous studies of this overall reaction sequence. While the thermal barrier for reversion of Q to N is relatively high, the energyreleasing conversion of Q to N can be achieved readily via a free-radical-cation chain reaction, initiated by oneelectron oxidations (Scheme 1).

Although the existence of two distinct radical cations,  $Q^{+}$  and  $N^{+}$ , had been established by chemically induced dynamic nuclear polarization (CIDNP) results,<sup>3</sup> the first direct observation of the quadricyclane radical cation ( $Q^{+}$ ) by time-resolved ESR (TR-ESR) was reported only recently.<sup>4</sup> A symmetric electronic structure was proposed for this elusive species and a detailed kinetic analysis indicated that its valence isomerization to  $N^{+}$  is relatively slow ( $k_3 = \sim 10^6 \text{ s}^{-1}$ ). A key question that remains unanswered in the dynamic interconversion of both the neutral Q–N system and their radical cations is their rates of interconversion.

An earlier theoretical study<sup>5</sup> provided geometries for Q and N that were in excellent agreement with electron diffraction experiments. Geometries were optimized at the HF/6-31G level and single point calculations which included d functions on carbon (6-31G\*) and second-order Møller-Plesset perturbation theory with the 6-31G basis were used to give estimated MP2 results with the 6-31G\* basis set. Despite the approximations involved in this study, it was clearly a state-of-the-art calculation at that time (121 basis functions). The estimated MP2/6-31G\* results for the energy difference between Q and N (26.7 kcal/mol) and Q++ and N++ (10.7 kcal/mol) were both only slightly higher than that observed experimentally (22  $\pm$ 1 and 9 kcal/mol).<sup>5,6</sup> Moreover, the estimates for the energy difference between the lower lying triplet state (T<sub>1</sub>) and N (60.7 kcal/mol) is in excellent agreement with the results presented below.

As part of a study<sup>7a</sup> on the potential involvement of  $Q^{\bullet+}$  in the cytochrome P-450 oxidation of  $Q^{,7b}$  we had occasion to optimize the geometry of both  $Q^{\bullet+}$  and  $N^{\bullet+}$ . We now report a theoretical prediction of the activation

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Scheme 1



barrier for the conversion of  $Q^{++}$  to  $N^{++}$  and the energy of the triplet state of Q relative to its ground state progenitor. Geometries have been optimized at the MP2/6-31G\* level while relative energies that are given have been calculated at the PMP4//MP2/6-31G\* and CCSD(T)//MP2/6-31G\* levels of theory.

At the PMP4//MP2/6-31G\* level of theory<sup>8</sup> we compute an energy difference between Q and N of 23.59 kcal/mol<sup>9</sup> (Table 1) in good agreement with the average of the experimental values (22  $\pm$  1 kcal/mol).<sup>5,6</sup> The triplet diradical state 2 is predicted to be 57.50 kcal/mol higher than N (Figure 1) in good agreement with an experimental values of 66.4 kcal/mol<sup>11a</sup> and 61.4 kcal/mol.<sup>11b</sup> Compared with the experimental geometries for N<sup>12</sup> and Q,<sup>13</sup> a modest improvement in the corresponding optimized geometries was observed over the HF/6-31G geometries<sup>5</sup> reported previously.<sup>14</sup> Triplet state **2** is slightly distorted to a lower symmetry  $(C_s)$  than its parent norbornadiene that has  $C_{2v}$  symmetry (Figure 2). The computed  $S^2$  for **2** at the MP2/6-31G\* level is 2.007, suggesting that this is a relatively pure spin state with little contamination from other multiplicities.

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<sup>(9)</sup> All geometries were fully optimized at the MP2/6-31G\* level of theory using the Gaussian 94 suite of programs<sup>10a</sup> with gradient optimization.<sup>10b</sup> Analytical frequency calculations on all stationary points were carried out at the MP2/6-31G\* level. Relative energies are given at the spin projected MP4 (PMP4) level<sup>8</sup> (including the ZPE correction unless otherwise stated) and the CCSD(T) level<sup>10c</sup> using MP2/6-31G\* geometries. All energies cited in the text are at the PMP4/6-31G\*/MP2/6-31G\* taking into account the zero-point energy unless specified otherwise.

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**Figure 1.** Photoisomerization of norbornadiene (1) to quadricyclane (3) and the thermal valence isomerization of quadricyclane radical cation **6** to norbornadiene radical cation **4** through transition state TS-**5**. Geometries are at the MP2/6-31G\*, and relative energies (kcal/mol) are at the MP4//MP2/6-31G\* and CCSD(T)//MP2/6-31G\* levels of theory. Experimental geometries are shown in parentheses.

 Table 1. Total Energies ar MP2/6-31G\*, MP4//MP2/6-31G\*, and CCSD(T)//MP2/6-31G\* Levels of Theory and Zero-Point Energies at the MP2/6-31G\* Level

molecule	MP2/6-31G*	MP4//MP2/6-31G*	PMP4//MP2/6-31G*	CCSD(T)//MP2/6-31G*	ZPE
norbornadiene 1 triplet min 2 quadricyclane 3 radical cation 4 transition state 5 radical cation 6	$\begin{array}{r} -270.553169\\ -270.458397\\ -270.520977\\ -270.265918\\ -270.230937\\ -270.256764\end{array}$	$\begin{array}{r} -270.644231\\ -270.549450\\ -270.607160\\ -270.356990\\ -270.322227\\ -270.343922\end{array}$	-270.551234 -270.357650 -270.325382 -270.345179	$\begin{array}{r} -270.644663 \\ -270.551859 \\ -270.606686 \\ -270.356474 \\ -270.323681 \\ -270.343376 \end{array}$	0.130565 0.129199 0.131086 0.134367 0.128421 0.130671

The calculated adiabatic ionization potential for conversion of N to N<sup>++</sup> is 7.90 eV and for Q to Q<sup>++</sup> is 7.12 eV in reasonable agreement with the experimental values of 8.37-8.43 and 7.40-7.86 eV, respectively.<sup>15</sup> The predicted  $S^2$  for N<sup>++</sup> of 0.7511 and for Q<sup>++</sup> of 0.7556 at the MP2/6-31G\* level are indicative of a relatively pure doublet state ( $S^2 = 0.75$ ) for both radical cations. The positive charge in Q++ and N++ is uniformly distributed between carbon atoms C<sub>2</sub>, C<sub>3</sub>, C<sub>5</sub>, and C<sub>6</sub>. The calculated atomic spin densities of  $C_2$  and  $C_6$  in the triplet (2) are 1.202 and the spin density of  $C_2$  in transition state  ${\bf 5}$  is 1.081. Transition state TS-5 for the valence isomerization of Q++ to N++ was fully optimized at the MP2/6-31G\* level of theory and has only one imaginary frequency (653i cm<sup>-1</sup>). The calculated activation barrier is predicted to be 11.01 kcal/mol at the PMP4/6-31G\*//MP2/6-31G\* level. A pulse radiolytic study of the Q-N system suggested an activation barrier of only  $20 \pm 5$  kJ/mol (4.8  $\pm$  1.2 kcal/mol).<sup>16</sup> An activation barrier of ~40 kJ/mol

(9.6 kcal/mol) has been suggested earlier based upon MINDO/3 calculations.<sup>13a</sup> This prompted us to do a single point calculation on TS-**5** at CCSD(T)/6-31G\*,<sup>10c,d</sup> which gave a barrier height that was only 0.6 kcal/mol lower in energy (Table 1). The predicted free energy of activation for isomerization of Q<sup>++</sup> to N<sup>++</sup> is  $\Delta G^{+}_{298} = 10.79$  kcal/mol. This value is considerably higher than the experimental barrier of 4.8 kcal/mol, obtained by Gebicki et al.<sup>16</sup> Transition state theory gives a calculated first-order rate constant  $k_3 = 0.76 \times 10^5$  s<sup>-1</sup> ( $A = 0.27 \times 10^{14}$  s<sup>-1</sup>,  $E_a = 11.66$  kcal/mol) in agreement with the more recent estimate of the experimental rate constant<sup>4</sup> of  $k_3 \sim 10^6$  s<sup>-1</sup>.

In summary, the radical cations derived from N and Q both have symmetrical electronic structures. The relative energies of both neutrals and radical cations are predicted in good agreement with experiment. The predicted half life for valence isomerization of Q<sup>•+</sup> to N<sup>•+</sup>,  $\tau_{1/2} = 9.14 \times 10^{-6}$  s at 298 K, is consistent with

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reaction coordinate

**Figure 2.** Relative energies (kcal/mol) at the PMP4//MP2/6-31G\* level of theory for the photoisomerization of norbornadiene (1) to quadricyclane (3) and the thermal valence isomerization of quadricyclane radical cation **6** to norbornadiene radical cation **4**. Values in parentheses are at the CCSD(T)//MP2/6-31G\* level.

experimental observations where the TR-ESR signal of  $Q^{\bullet+}$  decays within microseconds.

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