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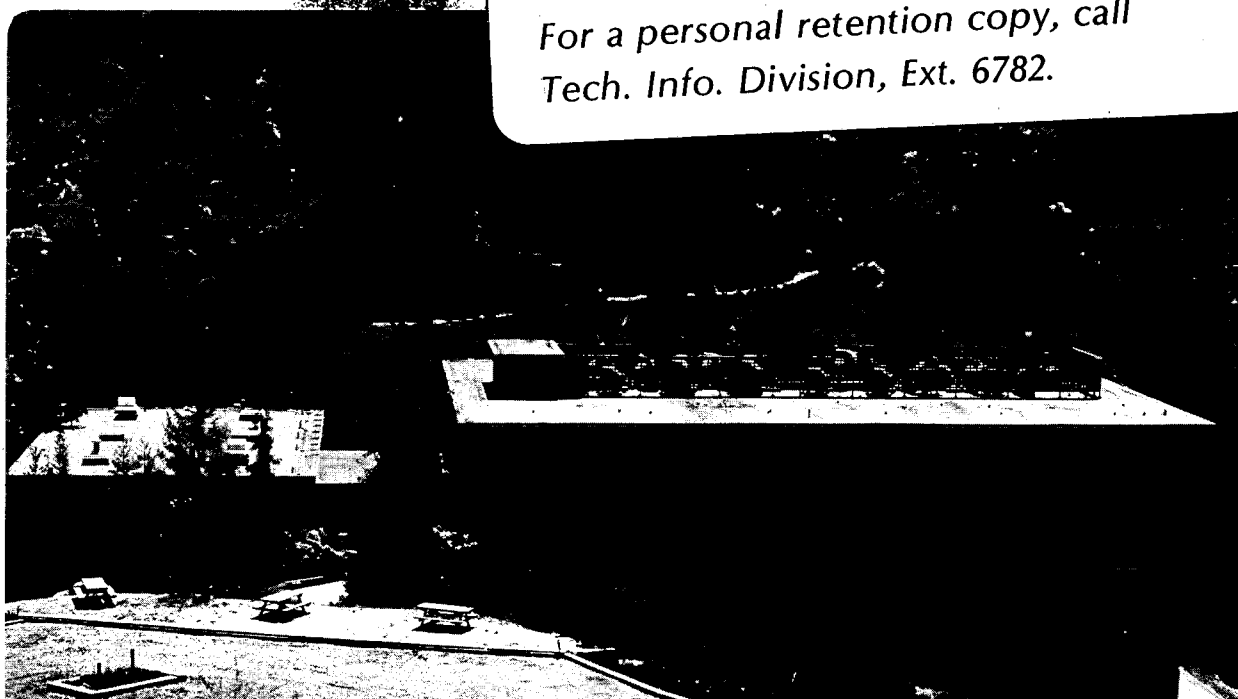
CYCLIC  $D_{6h}$  HEXAAZABENZENE--A RELATIVE MINIMUM ON  
THE  $N_6$  POTENTIAL ENERGY HYPERSURFACE?

Paul Saxe and Henry F. Schaefer III

July 1982

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Cyclic  $D_{6h}$  Hexaazabenzene--A Relative Minimum on the  $N_6$   
Potential Energy Hypersurface?

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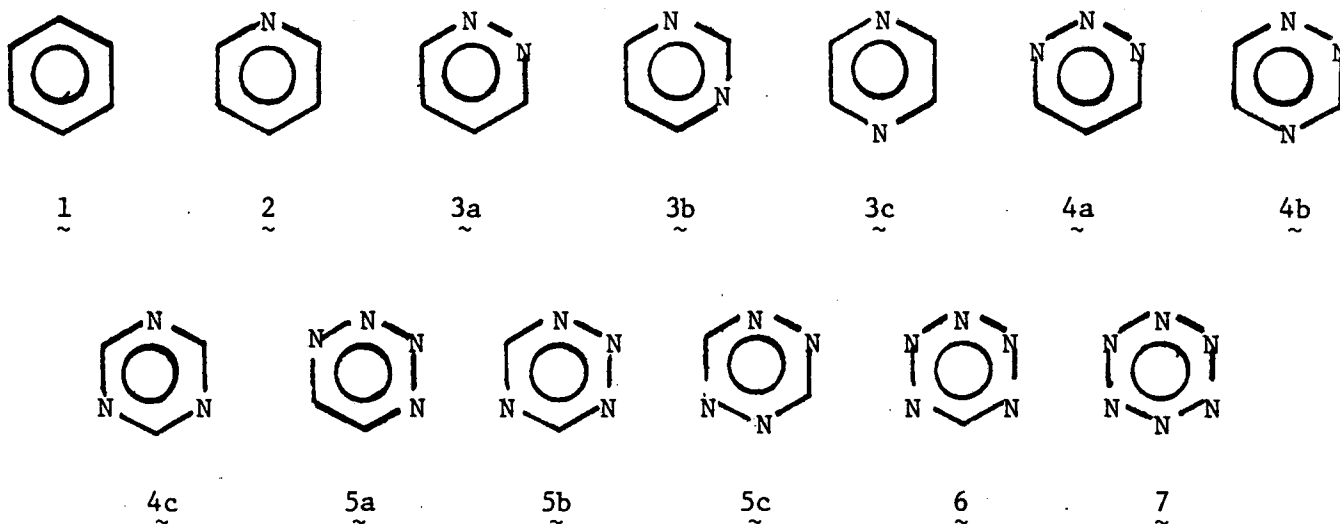
Abstract

Vogler has recently reported laboratory evidence for the formation of hexaazabenzene from photochemical elimination in cis-diazidobis(triphenylphosphorane) platinum(II). Previous theoretical studies have suggested that the  $D_{6h}$  benzene-like structure is not a minimum on the  $N_6$  potential energy hypersurface. Here the  $N_6$  problem has been addressed at the self-consistent-field (SCF) level of theory using double zeta (DZ) and double zeta plus polarization (DZ+P) basis sets. The smaller basis set yields the prediction that the  $D_{6h}$  structure is a transition state connecting two equivalent bond alternant  $N_6$  equilibrium geometries. A second transition state for dissociation to three nitrogen molecules (which are energetically much lower than  $N_6$ ) was also located. Contrary to previous theoretical work, hexaazabenzene is found to be a minimum at the highest completely consistent level of theory. The equilibrium geometry occurs for  $r_e(N-N) = 1.288 \overset{\circ}{\text{A}}$ , a bond distance suggesting that  $N_6$  is a classic aromatic molecule. The transition

state to  $3N_2$  lies 10.3 kcal higher and has the planar, bond alternant structure  $r_1(N-N) = 1.178 \text{ \AA}$ ,  $r_2(N-N) = 1.551 \text{ \AA}$ . Harmonic vibrational frequencies for hexaazabenzene are predicted with both theoretical methods and demonstrate that the energy surface is very flat with respect to bond alternating  $B_{2u}$  displacements. The inclusion of correlation effects lowers the barrier to  $N_6$  dissociation when geometrical structures obtained at the SCF level of theory are assumed.

Introduction

The  $N_6$  molecule hexazine or hexaazabenzene is the final member of the series of aromatic and potentially aromatic molecules

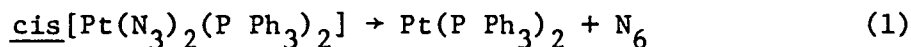


In this series of thirteen unsubstituted compounds, seven are well known and at least reasonably stable:<sup>1,2</sup> benzene (1), pyridine (2), pyridazine (3a), pyrimidine (3b), pyrazine (3c), s-triazine (4c), and s-tetrazine (5c). The eighth and last known compound of the series 1,2,4 triazine (4b), was synthesized in 1966<sup>3</sup> and an improved synthesis reported in 1974,<sup>4</sup> but 4b is apparently not well known, since a recent review<sup>1</sup> states that "the parent of this class has never been prepared."

Substituted examples of only one of the remaining five classes of compounds 1-7 have been prepared. The first 1,2,3 triazine (4a) prepared was the 4,5,6-triphenyl species reported by Chandross and Smolinsky<sup>5</sup> in 1960, and seventeen additional 1,2,3 triazines are

noted in Neunhoeffer's 1978 review.<sup>2</sup> Based on the stability of the 4,5,6-trimethyl-1,2,3 triazine, reported by Closs and Harrison<sup>6</sup> in 1972, it has been suggested<sup>2</sup> that the parent 4a should eventually prove (when synthesized) to be a relatively stable compound. However, there are no known examples of 1,2,3,4-tetrazines (parent 5a), 1,2,3,5-tetrazines (parent 5b) or pentazines (parent 6).

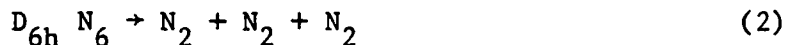
With this experimental background in mind, it is not at all surprising that no report of the synthesis of hexazine was reported prior to 1980. In a certain sense, of course, hexazine is the most intriguing of all the azabenzenes, since at least in principle it could share benzene's  $D_{6h}$  structure, all six CH fragments having been replaced by nitrogen atoms. For these reasons, the report by Vogler, Wright, and Kunkely<sup>7</sup> of evidence for the formation of hexaazabenzene was a startling one. These authors studied the low-temperature photolysis of the azide complex cis-[Pt(N<sub>3</sub>)<sub>2</sub>(PPh<sub>3</sub>)<sub>2</sub>] and analyzed the photoproducts in glasses of organic solvents (ethanol or methyltetrahydrofuran) at 77°K. A yellow coloration rapidly occurred (absorption maximum at  $\lambda = 380$  nm) which on warming the matrix immediately disappeared with simultaneous evolution of N<sub>2</sub>.<sup>7</sup> Vogler, Wright, and Kunkely (VWK) concluded that the N<sub>6</sub> molecule is formed as the azide ligands are cleaved, i.e.,



At room temperature, flash photolysis of the same platinum azide in the same solvents gave a similar uv absorption at  $\lambda = 380$  nm.

Prior to the VWK experiment,<sup>7</sup> there had already been at least six theoretical investigations<sup>8-13</sup> of hexazabenzene. As early as 1962, Roberts<sup>8</sup> noted that Hückel calculations of the electronic states of hexazabenzene would be the same as those for benzene, "provided the extra unshared electron pairs are regarded as being strictly localized." However, among quantitative theoretical studies, there is no compelling evidence that  $D_{6h}$  hexazine is a relative minimum on the  $N_6$  potential energy hypersurface. VWK appear to have been misled in this regard by certain statements in Wright's papers.<sup>11,12</sup> VWK state that Wright's calculations "lead to the conclusion that  $N_6$  is slightly stabilized," whereas in fact Wright only suggested<sup>12</sup> that  $D_{6h} N_6$  lies somewhat below  $N_2$  plus the highly energetic  $N_4$  species. Although  $N_6$  certainly does have "stability" in the sense attributed by Wright, such "stability" may have little bearing on whether  $N_6$  could be prepared in the laboratory. Since only  $D_{6h}$  geometries were considered in his work, Wright did not attempt to examine whether hexazabenzene was a relative minimum on the  $N_6$  potential surface.

In this light it may be noted that Dewar's MINDO/3 and MNDDO/1 studies<sup>13</sup> of the dissociation process



represent the earliest attempt to ascertain whether  $N_6$  might be an observable species in the gas phase. MINDO/3 predicts  $N_6$  to



be a relative minimum, while MNDDO/1 (stated to generally provide more satisfactory results) predicts that reaction (2) proceeds without an activation barrier. A more recent, and in some respects the most useful, study of the  $N_6$  problem is that of Ha, Cimiraglia, and Nguyen.<sup>14</sup> Using a 4-31G basis set<sup>15</sup> (of nearly double zeta quality<sup>16</sup>) they conclude (in concurrence with Dewar<sup>13</sup>) from self-consistent-field (SCF) and configuration interaction (CI) studies that even within  $D_{3h}$  symmetry, hexazine falls apart in the gas phase to  $3N_2$  without a barrier. This result notwithstanding, Ha<sup>14</sup> predicted an  $n \rightarrow \pi^*$  electronic transition at 391 nm, in reasonable agreement with VWK's observed<sup>7</sup>  $\lambda_{max} = 380\text{nm}$ . Ha concluded that harmony with the VWK experiment might be reached if hexazine were stabilized by the ethanol matrix at 77°K.

As this manuscript was in the final stages of preparation, a semi-popular review of theoretical aspects of the  $N_6$  problem appeared.<sup>17</sup> This review focussed our attention on the 1982  $N_6$  publication of Huber,<sup>18</sup> who applied the SCF method with a small double zeta basis including polarization by floating orbitals. Huber demonstrated via normal coordinate analysis that the  $D_{6h}$  form of  $N_6$  is not a minimum but a transition state at this level of theory. Huber states that "based on experience with calculations of this level, the result should not differ substantially by extending the basis". The present research indicates, among other things, that this suggestion by Huber is incorrect. That is, application of higher levels of theory to  $N_6$  does bring about qualitative changes in the potential energy hypersurface.

### Theoretical Perspective and Details

Thus the best theoretical calculations to date concur that hexaazabenzene is not a minimum on the  $N_6$  potential energy hypersurface. Although Ha<sup>14</sup> suggests the possibility that cyclic  $N_6$  would be matrix-stabilized, no information concerning the flatness or shape of the  $N_6$  energy surface near the optimum  $D_{6h}$  structure is provided. The present study picks up the  $N_6$  problem at this point and pursues it at a higher level of theory. Specifically, it may be mentioned that the existence of a minimum on the  $D_{6h}$  potential curve is of limited value. Since  $D_{6h} N_6$  has only one adjustable geometrical parameter, a minimum will necessarily exist as long as energies below that of six infinitely separated nitrogen atoms are found. Given the great strength of the  $N\equiv N$  bond in  $N_2$ , it is no surprise that all properly carried out theoretical studies to date agree that  $D_{6h} N_6$  lies well below  $6N$ . The goal of the present research was, then, to consider all the other vibrational degrees of freedom in the vicinity of the presumed hexaazabenzene structure in order to determine (a) whether the  $D_{6h}$  structure becomes a minimum at higher levels of theory and/or (b) whether the  $N_6$  energy hypersurface is sufficiently flat near the  $D_{6h}$  structure that stabilization by a matrix is plausible.

Two basis sets of contracted gaussian functions<sup>16</sup> were used in the present research. The first was the standard  $N(9s5p/4s2p)$  double zeta (DZ) basis of Huzinaga<sup>19</sup> and Dunning,<sup>20</sup> In the second basis, a set of six cartesian d-like functions (polarization

functions) with orbital exponent  $\alpha=0.8$  were added to each nitrogen atom. This double zeta plus polarization (DZ+P) basis set may be designated N(9s 5p 1d/4s 2p 1d). The DZ basis thus consists of 60 contracted gaussian functions and the DZ+P basis 96 contracted functions. All results presented here were obtained at the self-consistent-field (SCF) level of theory. It was possible to obtain configuration interaction (CI) wavefunctions and energies for a few selected geometries, but the SCF results (to be described below) indicated that such a procedure could be very misleading. That is, the  $N_6$  potential energy hypersurface is so flat near the constrained  $D_{6h}$  minimum that anything short of a complete geometrical optimization and harmonic vibrational analysis would appear to be of limited value.

Stationary point molecular geometries were determined using both analytic gradient<sup>21</sup> and direct energy search methods. The latter, although no longer generally helpful, proved useful in this context due to the extreme flatness of the  $N_6$  energy surface and the small number of independent geometrical parameters (1 for  $D_{6h}$  structures, 2 for  $D_{3h}$  structures). Given the  $D_{6h}$  stationary point structures, quadratic force constants were obtained as differences of analytic forces and subjected to standard harmonic vibrational analyses. DZ+P basis computation times on the Harris 800 minicomputer for SCF plus gradient were typically seven hours for geometries of  $C_{2v}$  symmetry and fifteen hours for point group  $C_s$ , with only a single reflection plane.

### Structures and Energetics

The DZ SCF and DZ+P SCF descriptions of  $N_6$  turned out to be very different, even in a qualitative sense. Therefore the lower level of theory will be discussed first. As seen in Figure 1, there are no less than three  $N_6$  stationary points in the vicinity of the anticipated  $D_{6h}$  hexazine structure. As will be demonstrated in the next section, the  $D_{6h}$  structure is indeed a transition state, as suggested by the nearly comparable 4-31G study of Ha, Cimiraglia, and Nguyen<sup>14</sup> and the work of Huber,<sup>18</sup> carried out with a small DZ basis set. Not mentioned by Ha or Huber is the fact that a very shallow (with respect to the  $D_{6h}$  transition state)  $D_{3h}$  minimum occurs nearby at the DZ SCF level of theory. This  $D_{3h}$  minimum lies only 0.05 kcal below the  $D_{6h}$  transition state and has unique bond distances  $r_1(N-N) = 1.295 \text{ \AA}$ ,  $r_2(N-N) = 1.353 \text{ \AA}$ , as seen in Figure 2. These distances correspond to an increase of  $0.031 \text{ \AA}$  in one bond distance and a compression of  $0.027 \text{ \AA}$  in the other, relative to the  $D_{6h}$  perfectly "aromatic" structure.

Since a  $D_{3h}$  equilibrium geometry was found by DZ SCF there must be a second transition state (in addition to the  $D_{6h}$  structure) connecting the  $N_6$  minimum with (the energetically much lower) three separated  $N_2$  molecules. This transition state is shown in Figures 1 and 2 and is quite well separated structurally from the other stationary points. Specifically, the longer N-N distance is increased by  $0.193 \text{ \AA}$  and the shorter distance decreased by  $0.098$  relative to the  $D_{6h}$  transition state. Despite this rather large shift in geometry, from two equal bond distances of  $1.322 \text{ \AA}$

to long and short N-N distances differing by  $0.291 \text{ \AA}$ , the accompanying energy difference is only 1.27 kcal. This finding suggests that the  $N_6$  energy difference is extraordinarily flat in the region of the  $D_{6h}$  hexaazabenzene structure. Although the DZ SCF characterization of the  $D_{6h}$  structure as a transition state will shortly be challenged, this more general conclusion of flatness is supported by the higher level results to be presented.

Contrary to previous theoretical studies,<sup>13,14,18</sup> the highest consistent level of theory yet used to address this problem suggests that  $D_{6h}$  hexazine is a relative minimum on the  $N_6$  energy surface. The fact that the DZ+P SCF method yields this result is demonstrated in Figure 1, with Figure 2 and Tables I and II providing additional details. The predicted N-N separation of the hexaazabenzene minimum is  $1.288 \text{ \AA}$ , or only  $0.034 \text{ \AA}$  shorter than the analogous DZ  $D_{6h}$  stationary point prediction. The striking difference, of course, is that with DZ SCF the  $D_{6h}$  structure is a transition state, while adding a set of d functions to each nitrogen atom transforms hexazine into a genuine potential minimum. Since the  $D_{6h}$  structure is a minimum with DZ+P SCF, there must be a transition state separating it from the dissociation limit of three nitrogen molecules. This transition state is seen in Figures 1 and 2, which show alternating bond distances of  $1.178$  and  $1.551 \text{ \AA}$ . Comparison of the DZ and DZ+P SCF results shows the surprising (relative to the structures of normal closed-shell molecules<sup>16</sup>) result that d functions increase the longer predicted N-N distance, by  $0.036 \text{ \AA}$ . The shorter N-N distance,

which is much more like that in isolated  $N_2$ , shows the expected decrease (by  $0.046 \text{ \AA}$ ) when polarization functions are added to the basis.

Quite unexpected relative to earlier theoretical work is the DZ+P SCF prediction (see Table I) that the  $D_{6h} N_6$  minimum is separated by a barrier of 10.3 kcal from the energetically much lower three  $N_2$  asymptote. Such a barrier is surprisingly large and, if correct, would readily explain Vogler's apparent low temperature observation<sup>7</sup> of  $N_6$ . Even if the precise value of the  $N_6$  dissociation barrier from DZ+P SCF theory is too high, the fact that the energy changes by so very little for such a broad range of N-N separations ( $1.551-1.178=0.373 \text{ \AA}$ ) is encouraging in terms of experimentally trapping  $N_6$  via matrix isolation techniques.

A matter of lesser interest here is the energy difference between  $D_{6h}$  hexaazabenzene and three separated  $N_2$  molecules. Previous theoretical studies<sup>10-14</sup> have demonstrated that hexazine lies energetically far above three  $N_2$  molecules. To make this comparison, the structure of the  $N_2$  molecule was optimized at the DZ SCF ( $r_e = 1.100 \text{ \AA}$ ) and DZ+P SCF ( $r_e = 1.083 \text{ \AA}$ ) levels of theory. The predicted bond distances are in acceptable agreement with the experimental value<sup>22</sup>  $r_e = 1.098 \text{ \AA}$ . Three times the SCF energies of  $N_2$  are given in Table I, from which a bit of subtraction shows that hexazine lies 231.7 kcal above  $3N_2$  with the DZ SCF method and 227.6 kcal above  $3N_2$  with DZ+P SCF. As Ha<sup>14</sup> has noted,

correlation effects will reduce these energy differences, inasmuch as  $D_{6h} N_6$  is described more poorly in the Hartree-Fock approximation than is the  $3N_2$  asymptote.

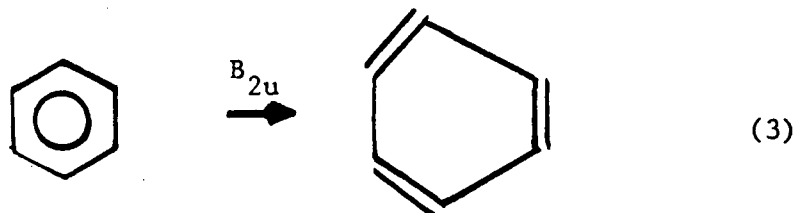
### Vibrational Frequencies of Hexaazabenzene

Consideration of the structure alone of  $D_{6h}$  hexazine would lead one to believe that it is, if anything, even more aromatic than benzene. Benzene has a C-C bond distance intermediate between typical C=C and C-C distances, being a bit closer to former. However, the predicted N-N distance in  $D_{6h}$  hexazine (1.322 Å DZ, 1.288 Å DZ+P) is very much closer to the double bonded diimide value<sup>23</sup> [ $r(N=N) = 1.25$  Å] than to that of the single bond in hydrazine<sup>24</sup> [ $r(N-N) = 1.447$  Å]. Furthermore, the predicted N-N distance is a bit shorter than the 1.32 Å found from the crystal structure<sup>25</sup> of the aromatic s-tetrazine molecule 5c. Thus the analogy between hexazine and benzene would appear to be a very good one based on the structural data presented here.

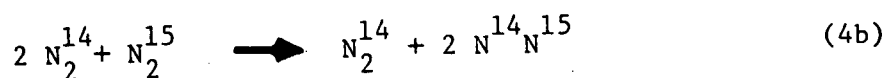
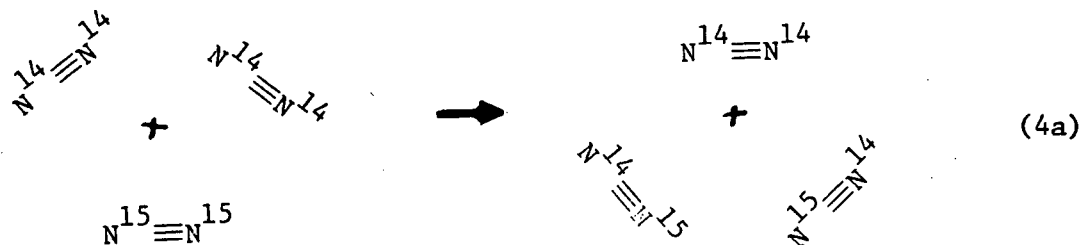
However, the predicted vibrational frequencies demonstrate clearly the dramatic differences between hexaazabenzene and benzene itself. These frequencies are given in Table II, where comparison is made with the benzene vibrational frequencies<sup>26</sup> associated with the carbon skeleton. Of course, some mixing between the C-H and C-C degrees of freedom is inevitable for benzene, so the comparison is not a perfect one.

From the DZ SCF results alone it is apparent that the  $B_{2u}$  vibrational frequencies of hexazine and benzene are about as

different as one could imagine. The  $B_{2u}$  mode allows one C-C distance to contract and the adjacent C-C bond to expand



destroying the aromaticity and moving toward a structure with distinguishable alternating single and double bonds. It has been known for years<sup>26</sup> that this is an expensive displacement energetically for benzene and this is reflected in the observed Kekulé stretching frequency of  $1309 \text{ cm}^{-1}$ . However for hexaazabenzene as modeled by DZ SCF, the  $B_{2u}$  mode becomes an imaginary vibration frequency (232 i), i.e., the  $D_{6h}$  structure is a maximum with respect to  $B_{2u}$  displacements. Thus the harmonic vibrational analysis of this species demonstrates conclusively that at the DZ SCF level of theory, hexazine is a transition state for nitrogen-nitrogen bond switching. One can imagine a degenerate isotopic reaction





that would employ such a transition state, but termolecular reactions of this general type are thought to be relatively unlikely.<sup>27</sup>

It is not without precedent for the nature of a stationary point to change in going from DZ to DZ+P SCF theory. For example, in the  $\text{SH}_4$  molecule, the  $\text{SF}_4$ -like  $\text{C}_{2v}$  minimum disappears when polarization functions are added to the basis set.<sup>28</sup>

Similarly for the  $\text{H}_7^+$  complex  $\text{H}_3^+(\text{H}_2)_2$  the known equilibrium geometry is predicted to be a transition state when CI is carried out with a DZ basis set.<sup>29</sup> In the same vein, the DZ SCF  $\text{D}_{6h}$  transition state becomes a minimum when d functions are added to the  $\text{N}_6$  basis set.

Of particular interest in Table II is the  $\text{B}_{2u}$  vibrational frequency predicted by DZ+P SCF. This frequency, while no longer imaginary (as with the DZ SCF method), is nevertheless very different from the experimental value for benzene. Specifically, Table II shows that the carbon skeletal  $\text{B}_{2u}$  frequency in  $\text{C}_6\text{H}_6$  is  $1309 \text{ cm}^{-1}$ , while the analogous DZ+P SCF frequency for hexazine is  $343 \text{ cm}^{-1}$ , almost a factor of four smaller. This is of course further concrete evidence that the  $\text{N}_6$  potential energy hypersurface is very flat in the region of the  $\text{D}_{6h}$  stationary point. The lowest DZ+P SCF vibrational frequency is not the  $\text{B}_{2u}$  mode but rather the degenerate  $\text{E}_{2u}$  frequency at  $320 \text{ cm}^{-1}$ . This is an out-of-plane bending mode quite analogous to that observed at  $404 \text{ cm}^{-1}$  for benzene. Thus one sees that the  $\text{N}_6$  energy hypersurface is only marginally flatter than benzene with respect to this ring deformation. In this general vein, one concludes that the DZ+P SCF

$N_6$  vibrational frequencies are all roughly comparable to the analogous skeletal modes of benzene except for the ring stretching  $B_{2u}$  Kekulé displacement.

#### Preliminary Results from Correlated Wave Functions

As noted earlier, it was not considered feasible at the present time to determine stationary point geometries and vibrational frequencies at the configuration interaction (CI) level of theory. However, assuming the SCF stationary point structures in Figure 2, single point CI wave functions were determined including all valence single and double excitations with respect to the  $N_6$  Hartree-Fock reference configuration. In  $C_{2v}$  point group (the symmetry of the SCF transition states for  $N_6 \rightarrow 3N_2$  is actually higher,  $D_{3h}$ ) there are a total of 33,320 configurations with the DZ basis set and 142,130 configurations with the DZ+P basis. These variational calculations were made possible by use of the recently developed shape-driven graphical unitary group approach,<sup>30</sup> and the larger CI required 12 hours on the Harris Corporation Series 800 minicomputer.

The results obtained from correlated wave functions are summarized in Table III. One sees first that electron correlation reduces the exothermicity of the  $N_6 \rightarrow 3N_2$  process by only  $\sim 17$  kcal, to  $\sim 211$  kcal. A much greater reduction ( $\sim 115$  kcal) in this exothermicity was earlier predicted by Ha, Cimiraglia, and Nguyen,<sup>14</sup> and we suspect that the latter result is not reliable, perhaps due to an improper use of configuration selection techniques.

With geometries determined at the SCF level of theory, the effect of electron correlation is apparently to raise the energy of the  $D_{6h}$  stationary point relative to adjacent regions of the  $N_6$  energy hypersurface. Thus with the larger DZ+P basis set the barrier to  $N_6$  dissociation is seemingly reduced to 1.3 kcal from the SCF value of 10.3 kcal. Moreover, appendage of the Davidson correction,<sup>31</sup> for unlinked clusters further reduces the energy of the geometry corresponding to the DZ+P SCF transition state, to 3.8 kcal below the energy of the analogous hexazine structure.

It is at best difficult to assess the significance of the correlated energy differences just reported. If the results obtained had involved complete geometry optimization at the correlated levels of theory, then one would be inclined to doubt the existence of hexazine as a relative minimum on the  $N_6$  potential surface. However, we must emphasize that no such correlated geometry determination was attempted and thus the results may be misleading. It should also be recalled that a counterintuitive result was found (in the longer of the alternating N-N distances for the  $N_6$  dissociation transition state) in going from the DZ to the DZ+P basis set. Thus it may not be possible to reliably estimate the effects of correlation of the  $N_6$  stationary point geometries. In conclusion, great progress has been made along the path to a reliable prediction of the properties of hexazine, but the process has not yet been carried to an entirely satisfactory conclusion.

### Concluding Remarks

The present theoretical study, carried out at a significantly higher level and degree of thoroughness than previous work, predicts the  $D_{6h}$  structure of hexaazabenzene to be a relative minimum on the  $N_6$  potential energy hypersurface. More pictorially, hexaazabenzene is predicted to be a dimple on top of a mountain. A question that must be addressed in this context is "Do the authors believe this result to be definitive?" To that question we must in candor give an equivocal answer. In the sense that we are confident that raising the level of theory applied will ultimately give correct answers to all such problems, then the DZ+P SCF prediction that  $N_6$  is a minimum must be considered more "reliable" than previous theoretical predictions to the contrary. However, this line of reasoning only suggests a most probable result and carries with it not a trace of infallibility. The primary and definitive conclusion of this research is rather that the  $N_6$  energy surface is very flat in the vicinity of the  $D_{6h}$  hexaazabenzene structure. Changes in the N-N bond distance of up to  $0.37 \text{ \AA}$  may be accommodated with a rise in the total energy of only 10 kcal.

Has hexaazabenzene actually been made<sup>7</sup> by Vogler, Wright, and Kunkely? At this point it appears to the present authors a bit premature to form an educated judgment. Further experiments are absolutely essential in this regard and it is hoped that the vibrational frequencies predicted here might assist in a matrix infrared spectroscopic identification of  $N_6$ . In any case, the unusual broadness of the  $N_6$  potential surface near the hexaazabenzene

stationary point does support the view that  $D_{6h} N_6$  might be trapped in a low-temperature matrix, as claimed by VWK.<sup>7</sup>

Acknowledgments

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Table I. Total energies (hartrees) and relative energies (kcal/mole) for  $N_6$  stationary points at various levels of theory. Relative energies are given with respect to the benzene-like  $D_{6h}$  stationary point. Note that while the  $D_{3h}$  structures reported here are definitely stationary points, they have not been proven by normal coordinate analysis to be minima and transition states, respectively.

	DZ SCF		DZ+P SCF	
	<u>Total Energy</u>	<u>Relative Energy</u>	<u>Total Energy</u>	<u>Relative Energy</u>
$D_{6h} N_6$	-326.26567 <sup>a</sup>	0.00	-326.51427	0.00
$D_{3h} N_6$ minimum	-326.26574	-0.05		No such stationary point
$D_{3h} N_6$ transition state	-326.26364	+1.27	-326.49784	+10.3
Three $N_2$ molecules	-326.63493	-231.7	-326.87698	-227.6

<sup>a</sup>An energy of -326.26557 hartrees (0.0001 hartree higher) was reported by Ha, Cimraglia, and Nguyen<sup>14</sup> apparently using the identical DZ SCF method. Their  $D_{6h}$  geometry is  $r(N-N) = 1.320 \text{ \AA}$ , while the present stationary point geometry is  $r(N-N) = 2.4974 \text{ bohrs} = 1.322 \text{ \AA}$ .

Table II. Predicted vibrational frequencies (in  $\text{cm}^{-1}$ ) for hexaazabenzene compared with those known experimentally for benzene. To provide a more realistic comparison, the DZ+P SCF harmonic frequencies have also been reduced by 10% and these empirically corrected frequencies are given in parentheses. Experimental frequencies and assignments are from reference 26.

<u>Normal Mode</u>	<u>Hexazine DZ SCF</u>	<u>Hexazine DZ+P SCF</u>	<u>Benzene Experimental</u>	<u>Benzene Assignments</u>
In-plane frequencies				
$E_{2g}$	1539	1723 (1551)	1599	C-C stretching
$E_{1u}$	1257	1412 (1271)	1482	C-C stretching
$B_{1u}$	1160	1258 (1132)	1010	C-C-C trigonal bending
$A_{1g}$	1083	1185 (1067)	993	breathing
$E_{2g}$	769	809 (728)	606	C-C-C in-plane bending
$B_{2u}$	2321	343 (309)	1309	C-C stretching (Kekulé)
Out-of-plane frequencies				
$B_{2g}$	913	959 (863)	707	C-C-C puckering
$E_{2u}$	288	320 (288)	404	C-C-C out-of-plane bending

Table III. Total energies (in hartrees) and relative energies (in kcal/mole) from correlated wave functions for  $N_6$ . Relative energies are given with respect to the benzene-like  $D_{6h}$  stationary point. Note that the geometries used here are those in Figure 2, i.e., optimized at the single configuration SCF level of theory. Thus reoptimizing the structures (assumed here) at the CI level could significantly effect these predictions.

	DZ CI		DZ+P CI	
	<u>Total Energy</u>	<u>Relative Energy</u>	<u>Total Energy</u>	<u>Relative Energy</u>
$D_{6h} N_6$	-326.80802	0.0	-327.28247	0.0
$D_{3h} N_6$ minimum	-326.80884	-0.5	--	--
$D_{3h} N_6$ transition state	-326.82266	-9.2	-327.28035	+1.3
Three $N_2$ molecules	-327.17498	-230.3	-327.61638	-209.5

Figure Captions

- Figure 1. Qualitative sketches of the potential energy surfaces connecting  $D_{6h}$  hexaazabenzene with three infinitely separated  $N_2$  molecules. As a one-dimensional reaction coordinate, the longer of the two distinct N-N distances in  $D_{3h}$   $N_6$  was chosen. Of course the two N-N distances coalesce as one passes from  $D_{3h}$  to  $D_{6h}$  point group.
- Figure 2. Stationary point geometrical structures for  $N_6$ . N-N bond distances are given in Å. Note in Figure 1 that the DZ SCF and DZ+P SCF potential surfaces are very different.

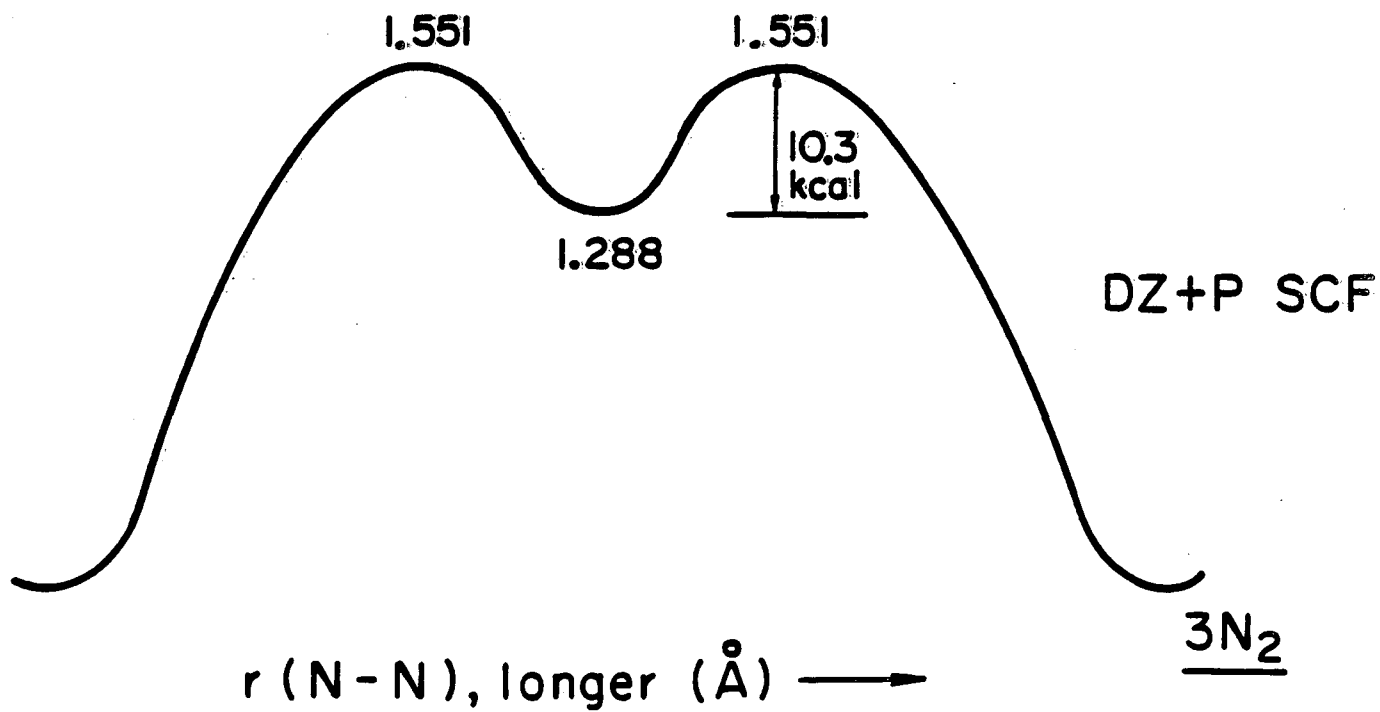
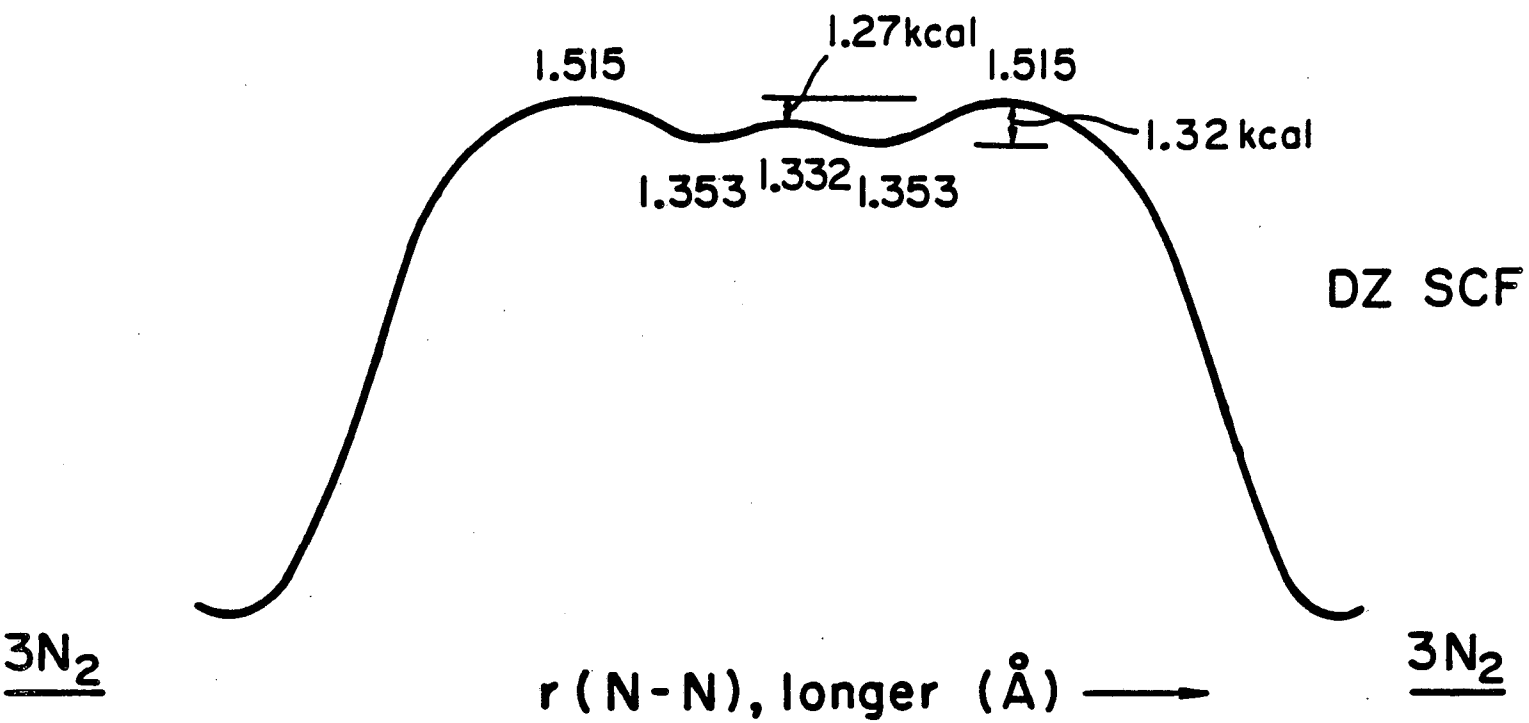


Figure 1

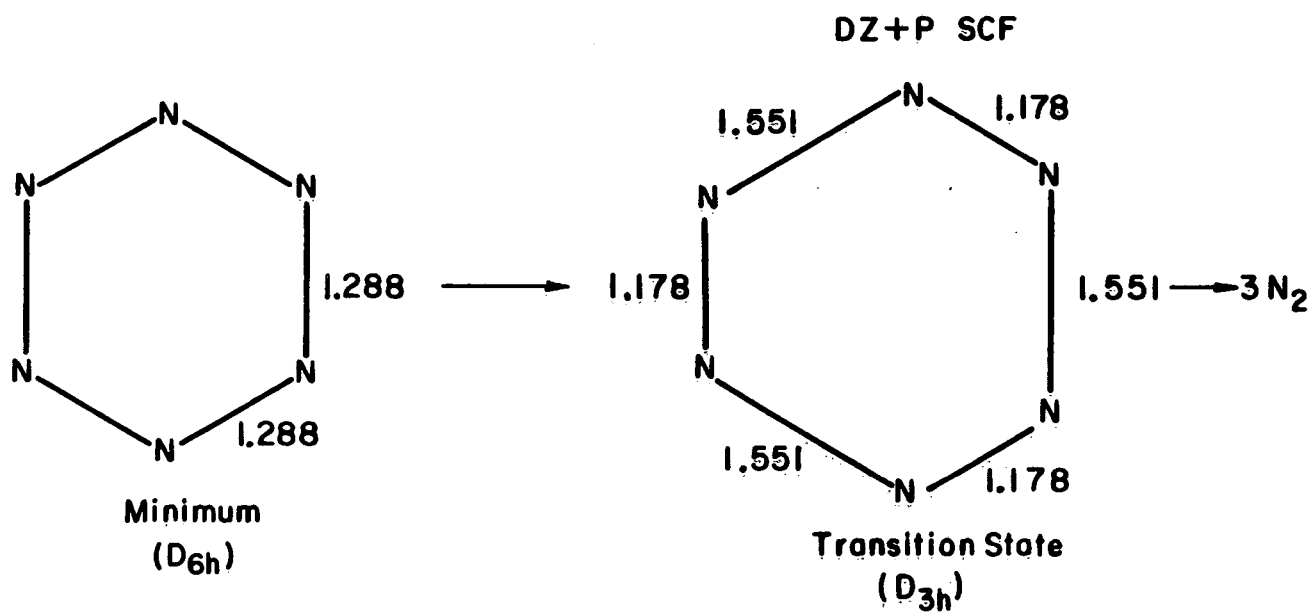
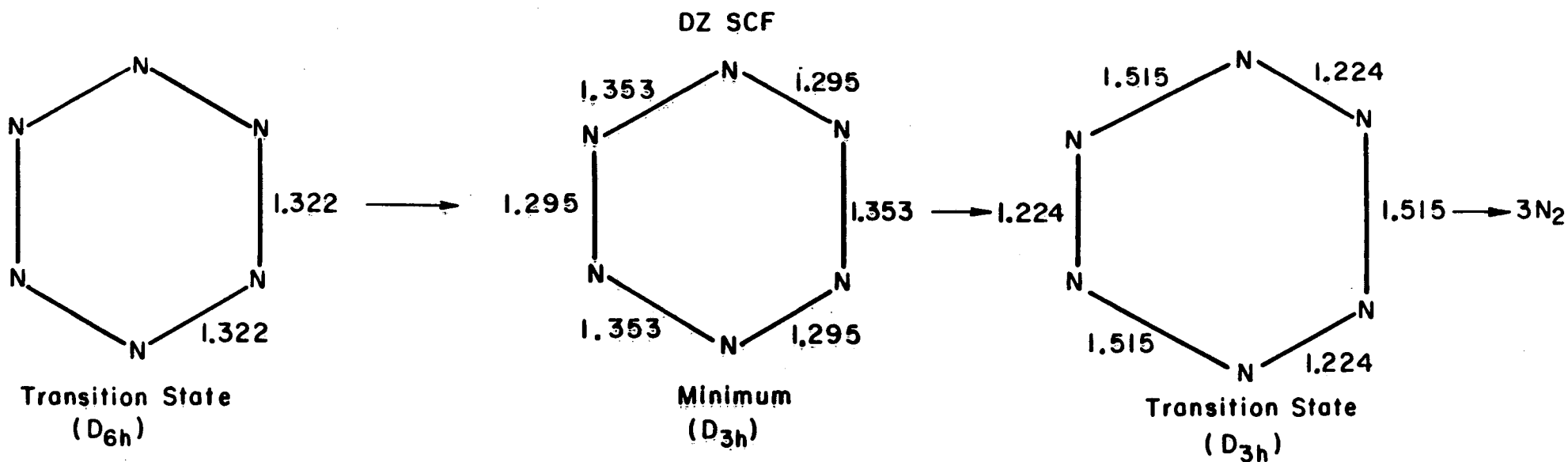


Figure 2

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