Photolysis Procedures. The light source most frequently used was a 400-W, medium-pressure, mercury, General Electric lamp H400 A-33-1. However, removal of the outer glass shell from this street lamp and slight bending of the electrical connections make it suitable as an inexpensive source of ultraviolet radiation. Light in the vicinity of 313 nm was isolated by placing the mercury lamp in a water-cooled quartz well filled with an aqueous solution, 0.0025 M in K<sub>2</sub>CrO<sub>4</sub> and 1% in K<sub>2</sub>CO<sub>3</sub>.<sup>27</sup> The jacket surrounded the lamp with a 7-nm thickness of filter solution.

The solutions to be photolyzed were placed in 75-mm long, cylindrical, 20 mm o.d. Pyrex tubes. The tops of the tubes were fitted with 14/20 ST outer joints. The tubes could be sealed with a 14/20 ST inner joint to which was attached a Pyrex ST stopcock.

The contents of the tubes were first frozen with either liquid nitrogen or a dry ice-acetone slurry. The stopcock to the tube was then opened, and the tube was evacuated to 0.005 Torr. The stopcock was closed and the contents of the tube was allowed to thaw. The cycle was repeated four times.

To ensure that the samples received equal amounts of incident light during the quantum yield determinations, the tubes to be photolyzed were rotated about the light source at 5 revolutions/ min with a Southern New England Ultraviolet Company Model MGR 500 "merry-go-round".

The samples to be irradiated for quantum yield determinations were maintained at 30  $\pm$  0.4° throughout the photolysis by immersing the samples, "merry-go-round", and light source into a constant-temperature bath.

For product studies, samples were photolyzed in degassed Pyrex tubes, suspended in a Rayonet photochemical reactor, equipped with 300-nm bulbs. The light from the bulbs was filtered by suspending the solutions in the same dichromate carbonate solution used as a filter in quantum yield determinations and quencher studies.

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# Photoelectron Spectra and Molecular Properties. L.<sup>1-3</sup> On the Blue Color of Bis(trimethylsilyl)diimine

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Abstract: Photoelectron (PE) and uv spectra of group 4 azo compounds  $R_3X - N = N - YR_3$  (X, Y = C, Si) and  $H_5C_6 - N = N - YR_3$  (X, Y = C, Si) and  $H_5C_6 - N = N - YR_3$  (X, Y = C, Si)  $N=N-XR_3$  (X = C, Si, Ge) are reported. A comparative discussion, based on CNDO calculations, attributes the low ionization potentials of the trimethylsilyl derivatives to substantial electron donation from the R<sub>3</sub>Si substituents. The unique electronic spectrum of the title compound can be rationalized considering in addition electron-electron interaction as well as Rydberg excitations.

It is the blue color of bis(trimethylsilyl)diimine which has stimulated investigation of its molecular properties<sup>3,4</sup> as well as those of related compounds.<sup>2,3,5</sup> In spectroscopic terms, its long wavelength  $n_N^+ \rightarrow \pi^*$  transition<sup>6-9</sup> is batho-

chromically shifted by more than 14000  $cm^{-1}$  (!) relative to that of the isoelectronic tert-butyl derivative.<sup>3,6,7</sup> As pointed out for the corresponding ketones,<sup>5</sup> analogous substituent dependent shifts of the long wavelength absorption are also observed for other chromophoric systems. For bis(trimethylsilyl)diimine and its challenging spectroscopic prop-

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erties tentative explanations range from inductive destabilization of the nitrogen electron pairs and/or silicon acceptor orbital participation<sup>2-6</sup> to a widening of the angle NNSi.<sup>3</sup> In the meantime the question of  $d_{\pi}p_{\pi}$  bonding in organometalloids is widely accepted to be a rather semantic one<sup>10-12</sup> and a recent x ray structure determination proved an angle NNSi of 120°.<sup>13</sup> Although other silicon nitrogen com-



pounds exhibit a comparable angle<sup>13</sup> of ~120°, it is increased relative to angles CNN ~ 106-113° found for unstrained azo alkanes. The most striking difference, however, is the rather short NN bond length  $d_{\rm NN} = 1.17$  Å,<sup>13</sup> halfway between values for carbon azo compounds with  $d_{\rm NN} = 1.23-1.27$  Å and the nitrogen molecule with  $d_{\rm NN} = 1.11$  Å. This finding can be attributed to an unusual high electron density in the NN bond of bis(trimethylsilyl)diimine and already suggests a considerable electron donation from the adjacent silicon substituents.

#### **PE Spectra and Assignment**

In order to support the remaining explanation for the blue color of bis(trimethylsilyl)diimine, i.e., a model emphasizing inductive effects, PE spectra of some group 4 azo compounds were recorded (Figure 1 and Table I).

The first PE band—well separated in the spectra of most azo compounds<sup>7,14</sup>—is readily assigned to ionization from the nitrogen electron pairs (Figure 2:  $n_N^+$ ). This can be substantiated not only by the results of CNDO calculations but also by comparison with the carefully analyzed PE spectrum of azomethane<sup>14</sup> and by general uv spectroscopic arguments.<sup>6-9</sup> As expected,<sup>15</sup> the corresponding ionization energy, IE<sub>1</sub>, is lowered on replacing *tert*-butyl by trimethylsilyl or trimethylgermyl substituents (Table I). Thereby a nearly constant shift  $\Delta IE_1$  occurs on stepwise substitution for the diimine derivatives  $R_3X$ —N=N—XR<sub>3</sub>. This observation favors the application of an inductively dominated model, in agreement with, e.g., the CNDO charge densities at the nitrogen atoms  $q_N$  of the neutral molecules (values







Figure 1. PE spectra of bis(trimethylsilyl)diimine, bis(tert-butyl)diimine, and of corresponding phenyl azo derivatives.

**Table I.** Vertical Ionization Potentials ( $IE_n$ , eV) of Bis(trimethylsilyl)diimine, 1-Trimethylsilyl-2-tert-butyldiimine, Bis(tert-butyl)diimine, and of Some Group 4 Phenylazo Compounds,  $C_6H_s$ —N=N—X( $CH_3$ )<sub>3</sub>

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	x	Y	IE,	IE2	IE3	IE4	IE,
YR,	Si	Si	7.1	10.6	12.45	13.5	
R <sub>3</sub> X N=N	Si C	C C	7.6 8.2	10.4 10.9	11.0 11.5	12.6	
N N		С	8.35	9.1	9.35	11.1	12.5
		Si Ge	7.85 7.65	9.1 9.0	9.4 9.35	10.7 $10.2$	11.8 11.6
*							

ditional CNDO calculations including  $3d_{Si}$  orbitals into the basis set did neither significantly improve the results nor provide new insight. In accordance with previous PE spectroscopic assignments based on CNDO calculations,<sup>11</sup> the conclusion therefore remains that there seems to be no way to definitely prove or disprove  $3d_{Si}$  orbital participation.<sup>10-12</sup>

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Very little else can be assigned in the rather featureless diimine PE spectra (Figure 1); even the number of ionizations hidden under the humps cannot be extracted. This is due to the numerous  $\sigma$ -type ionizations in the 28-atom molecules (H<sub>3</sub>C)<sub>3</sub>X—N=N-X(CH<sub>3</sub>)<sub>3</sub> containing 60 valence electrons. In general, three characteristic molecular orbitals are expected for the azo group: the nitrogen electron pair combinations n<sub>N</sub><sup>+</sup> and n<sub>N</sub><sup>-</sup> and the  $\pi$  bonding orbital  $\pi_{NN}$ . The relative energies of these orbitals have been discussed<sup>14</sup> in terms of a through space splitting<sup>16</sup> into n<sub>N</sub><sup>-</sup>/n<sub>N</sub><sup>+</sup> with a subsequent through bond destabilization<sup>15</sup> shifting the n<sub>N</sub><sup>+</sup> orbital above n<sub>N</sub><sup>-</sup>. The lowest ionization potential therefore is assigned to the n<sub>N</sub><sup>+</sup> orbital combination (Figures 1 and 2).

However, semiempirical MO calculations even on simple azo compounds such as azomethane suggest that the classification given in Figure 2 represents an unrealistically oversimplified picture.<sup>14</sup> All the more, for the complicated molecules dealt with in this paper any such classification hardly seems possible. Judging from the CNDO calculation only the  $n_N^+$  and  $\pi_{NN}$  orbitals can still be recognized, whereas  $n_N^-$  is heavily mixed with numerous  $\sigma$ -bonding orbitals of the substituents X(CH<sub>3</sub>)<sub>3</sub>. This is illustrated by, e.g., a sketch of the disilyldiimine molecular orbitals  $\psi_{24}$  (a) and  $\psi_{23}$  (b) with highest  $n_N^-$  and  $\pi_{NN}$  contributions, respectively:



For example, in  $\psi_{23}(n_N^-)$  the nitrogen orbital components amount to only one-third of the total, and thus the interesting problem of substituent dependent magnitude of the  $n_N^+/n_N^-$ -splitting finally must be abandoned because no (canonical) molecular orbital can be identified as  $n_N^-$  in more complex azo compounds.

To end the PE spectroscopic discussion of the diimine derivatives  $R_3X - N = N - XR_3$ , in Figure 3 the detectable lower ionization potentials (Table I) are compared with the CNDO eigenvalues assuming validity<sup>17</sup> of Koopmans theorem IE<sub>n</sub> =  $-\epsilon_J^{SCF}$ .

In view of the complexity of the molecules dealt with, the overall PE spectroscopic pattern is satisfactorily reproduced, e.g., the 3-eV gap between the first two ionizations. According to the calculations, some  $\sigma$  orbitals are inserted



Figure 2. Qualitative splitting scheme for the characteristic molecular orbitals of an azo group and the first three ionization energies of azomethane.<sup>12</sup>



Figure 3. Comparison of ionization energies  $IE_n$  (eV) and CNDO eigenvalues  $\varepsilon_J^{CNDO}$  (eV).

between  $n_N^+$  and  $\pi_{NN}$ , which also seems reasonable with respect to the PE spectra of tetramethylsilane<sup>18</sup> and 2,2dimethylpropane.<sup>19</sup> However, no evidence can be gathered from the PE spectra as to where to locate the  $\pi_{NN}$  ionizations. An approximate estimate based on the  $IE_n/-\epsilon_J^{SCF}$ correlation (Figure 3) yields for bis(trimethylsilyl)diimine a difference  $IE(\pi_{NN}) - IE(n_N^+)$  of more than 3 eV. For the PE spectra of the phenyl azo derivatives

For the PE spectra of the phenyl azo derivatives  $C_6H_5$ —N=N—XR<sub>3</sub> (Figure 1 and Table I) a perturbation approach suggests that the degeneracy of  $e_{2g}$  benzene orbitals (IE<sub>1</sub>(C<sub>6</sub>H<sub>6</sub>) = 9.24 eV<sup>15</sup>) is removed by inductive and conjugative effects of the azo group. The shift of the first PE band again reflects increasing (inductive + hyperconjugative) electron donation from the substituents CR<sub>3</sub> < SiR<sub>3</sub> < GeR<sub>3</sub>.

Summarizing, the PE spectra of the group 4 azo compounds are rather poorly structured and thus—besides some phenyl-type ionizations—only the strongly substituent dependent first bands can be unambiguously assigned. Nevertheless, based on the energies of the  $n_N^+$  orbital combinations—established above within the limits and implications of Koopmans' theorem<sup>17</sup>—some comments on the optical absorption spectra and especially on the electronic transitions of bis(trimethylsilyl)diimine are rendered possible.

#### Absorption Spectra

The optical absorption spectra (Figure 4) have been recorded in hydrocarbon solvents except for the gaseous phase

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Figure 4. Optical absorption spectra of some group 4 azo compounds  $R_3X - N = N - XR_3$  and  $H_5C_6 - N = N - XR_3$ .<sup>20</sup>

spectrum of bis(*tert*-butyl)diimine. Transition energies and molar extinction coefficients are given in Table II.

In all electronic spectra<sup>20</sup> the  $n^+ \rightarrow \pi^*$  bands are wellseparated from the high intensity absorption regions, displaying a characteristic long wavelength shift on  $R_3C \rightarrow$  $R_3Si$  substituent exchange. The  $n^+ \rightarrow \pi^*$  excitation energies of the  $R_3Si$  and  $R_3Ge$  derivatives differ only little. The uv region of the phenylazo compounds is dominated by transitions from the aromatic system<sup>21</sup>—also in accord with the assignment of the second and third ionization potentials (Table I) to predominantly phenyl-type molecular orbitals. In contrast, the  $R_3X-N=N-XR_3$  derivatives show intense absorptions only near 50000 cm<sup>-1</sup>, and again bis(trimethylsilyl)diimine exhibits a special feature. Besides the lowest  $n_N^+ \rightarrow \pi^*$  excitation energy observed so far for an azo compound at the borderline between the visible and infrared spectral regions, there appears a second band of moderate intensity at about 40000 cm<sup>-1</sup> (Figure 4 and Table II).

The electronic spectra even of simple azo compounds are still puzzling; in general, only the  $n_N^+ \rightarrow \pi^*$  transition can be unequivocally assigned, whereas, e.g., the location of the  $\pi \rightarrow \pi^*$  band has been disputed again and again.<sup>6-9,22</sup> For a discussion of the bis(trimethylsilyl)diimine excitations we start with an approach applied to azomethane by Haselbach and Schmelzer.<sup>8</sup> On the same level of approximation as Koopmans' theorem,<sup>17</sup> the electronic excitation energies are given by<sup>23</sup>

$$\Delta E_{\mathbf{i} \rightarrow \mathbf{k}} = \epsilon_{\mathbf{k}} - \epsilon_{\mathbf{i}} - J_{\mathbf{i}\mathbf{k}} + 2K_{\mathbf{i}\mathbf{k}}$$

correcting the orbital energy difference  $\epsilon_k - \epsilon_i$  for changes in interelectronic repulsions, i.e., by coulomb integral  $J_{ik}$ and by exchange integral  $K_{ik}$  terms. Within the CNDO approximation and parametrization,<sup>23</sup> values of

$$J_{ik} = \int \int \psi_i^{+}(1)\psi_k^{+}(2)r_{12}^{-1}\psi_i(1)\psi_k(2) d\tau_1 d\tau_2$$
  
$$K_{ik} = \int \int \psi_i^{+}(1)\psi_k^{+}(2)r_{12}^{-1}\psi_k(1)\psi_i(2) d\tau_1 d\tau_2$$

have been calculated for the  $n \rightarrow \pi^*$  and  $\pi \rightarrow \pi^*$  transitions of bis(trimethylsilyl)dimine.

ransition	$J_{ik}$	K <sub>ik</sub>	$J_{ik} = 2K_{ik}$	$\Delta E_{i \rightarrow k}$	
$n_N^{+} \xrightarrow{\pi^*} \pi^*$	8.65	0.00	8.6	0.9	(eV)
$\pi \xrightarrow{\pi^*} \pi^*$	8.65	1.36	5.9	10.5	(eV)

Although no emphasis is put on the actual numerical values, obviously the  $K_{ik}$  terms differ considerably for the different electronic transitions of the disilyl azo compound. According to ref 8, small  $K_{ik}$  values can be traced back to only minor contributions of same basis functions to the two molecular orbitals under consideration, whereas small Jik values indicate considerable orbital extension outside the region of the azo group.<sup>8</sup> All in all, reasoning along these lines Hasselbach and Schmelzer<sup>8</sup> have accounted for the fact that the difference between  $n_N^+$  and  $\pi$  ionization potentials in azomethane is smaller than the difference between  $n_N^+ \rightarrow \pi^*$  and  $\pi \rightarrow \pi^*$  excitation energies. For bis-(trimethylsilyl)diimine a difference in  $J_{ik} - 2K_{ik}$  of 2.7 eV is calculated, suggesting a much larger distance between n  $\rightarrow \pi^*$  and  $\pi \rightarrow \pi^*$  bands in the electronic spectrum than expected from the ionization potential difference IE  $(\pi_{NN}) - IE(n_N^+)$  alone. Finally, the CNDO excitation energy  $\Delta E_{\pi \to \pi^*}$ , even though its numerical accuracy might be questioned, would place the  $\pi \rightarrow \pi^*$  band in the vacuum uv region at about 80000 cm<sup>-1</sup>.

As pointed out before, the assignment of the  $\pi \to \pi^*$ transition in the electronic spectra of azo compounds R-N=N-R is not yet settled.<sup>8,9,22</sup> In azoalkanes the  $\pi \to \pi^*$ excitation energy possibly exceeds 70000 cm<sup>-1,9,22</sup> For the observed transitions in between  $n_N^+ \to \pi^*$  and  $\pi \to \pi^*$  two

Table II. Transition Energies ( $\nu_m$ , cm<sup>-1</sup>) and Molar Extinction Coefficients ( $\epsilon_m$ , 1./(mol cm)) of Bis(trimethylsilyl)diimine,<sup>b</sup> 1-Trimethylsilyl-2-tert-butyldiimine,<sup>b</sup> Bis(tert-butyl)diimine, and of Phenylazo Compounds, C<sub>6</sub>H<sub>5</sub>-N=N-X(CH<sub>3</sub>)<sub>3</sub><sup>b</sup> with X = C, Si, Ge<sup>20</sup>

	x	Y	$\widetilde{\nu}_{_{1}}/\epsilon_{_{1}}$	$\widetilde{\nu}_2/\epsilon_2$	$\widetilde{\nu}_{3}/\epsilon_{3}^{a}$	$\widetilde{\nu}_{4}/\epsilon_{4}^{a}$
YR	Si	Si <sup>b</sup>	12750/5	40000/260	52100/1660	57000/2680
N=N	Si	C <sup>b</sup>	20000/9	50000	)/1080	
R <sub>a</sub> X	С	C <sup>c</sup>	27200/12	50000/1800		59500/6800
XR	С	Ь	24400/144	38400/8400	47600/11900	
N=N	Si	b	17250/29	37500/16350	40000/14750	45500/15400
Í						46600/18100
$\bigcirc$	Ge	Ь	17800/26	39200/6780		46800/7840

<sup>a</sup> Maxima of undeconvoluted bands. <sup>b</sup> Solvent: hydrocarbon (pentane, hexane, or heptane). <sup>c</sup> Gas phase.

assignments are discussed in the literature:  $n_N \rightarrow \pi^{*6,8,22}$ and  $n_N^+ \rightarrow np$  Rydberg.<sup>9</sup> The latter proposal by Robin<sup>9</sup> is based on the following general procedure.<sup>24</sup> The Rydberg term value  $R/(n-\delta)^2$  is obtained by subtracting the uv excitation energy  $hv_1$  from the photoelectron spectroscopic ionization potential IE<sub>1</sub>. The quantum defect  $\delta$  calculated from the term value allows one to predict the excitation energies  $hv_{1+m}$  of higher members n + m (m = 1, 2, 3...)



of the corresponding Rydberg series.<sup>24</sup> For azomethane, the  $n_N^+ \rightarrow 3p$  Rydberg assignment of the band at 54000 cm<sup>-1</sup> is not only supported by the correct prediction  $\nu(n_N^+ \rightarrow 4p)$ Rydberg)  $\sim$ 63000-64000 cm<sup>-1</sup> of the next band but also by the vanishing vibrational fine structure in the spectrum recorded under 119 atm of nitrogen pressure.9

Comparison of the azomethane values with those for the symmetrically disubstituted azo compounds R-N=N-R shows term values IE<sub>1</sub> -  $\nu_1$ , quantum defects  $\delta$ , and band distances  $\nu_2 - \nu_1$  of the same order of magnitude (Table III). In addition, the  $n_N^+ \rightarrow 5p$  Rydberg excitation is predicted analogously at about 52000  $cm^{-1}$  (cf. Figure 4). The satisfying fit of both the presumably 4p and 5p Rydberg transitions to the recorded spectrum lends some credit to a tentative assignment  $n_N^+ \rightarrow 3p$  Rydberg for the long wavelength band at 40000 cm<sup>-1</sup> of bis(trimethylsilyl)diimine. Its rather low intensity ( $\epsilon_{mol} \sim 260 \text{ l.}/(\text{mol cm})$ ) is in accord with the uv spectra of azo alkanes which show for the corresponding band a considerable drop of the molar extinction with increasing size of the alkyl group, e.g., for H<sub>3</sub>C- 7500, and for (H<sub>3</sub>C)<sub>2</sub>HCCH<sub>2</sub>- 1200 1./(mol cm).<sup>9</sup> This observation can probably be explained by a distortion of the symmetry of lower Rydberg states accompanied by intensity redistribution.9

#### **Concluding Remarks**

Summarizing facts and speculations, the electronic spectrum of the blue title compound, bis(trimethylsilyl)diimine, is unique not only because of the tremendous  $n_N^+ \rightarrow \pi^*$ shift relative to azo alkanes but also due to the strong electron donation by R<sub>3</sub>Si substituents. In addition, there is some evidence for a low-lying 3p Rydberg state.

Rydberg states of polyatomic molecules, as defined in ref 24, can play a rôle in photochemistry.<sup>25</sup> For example, Sakurai and co-workers<sup>26</sup> reported that 1,2,3-trisilacycloheptanes expel almost quantitatively dialkylsilylenes when irradiated with a low pressure mercury arc lamp. Ramsey<sup>27</sup> has suggested a three-membered ring transition state for this reaction. The first ionization potential of octamethyltrisilane<sup>28</sup> amounts to IE<sub>1</sub> = 8.19 eV  $\simeq$  66000 cm<sup>-1</sup>, and the

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R	CH <sub>3</sub>	C(CH <sub>3</sub> ) <sub>3</sub>	Si(CH <sub>3</sub> ) <sub>3</sub>	
IE,	72400	66100	57300	cm <sup>-1</sup>
ν,	54200	49000a	40000	cm -1
$I\dot{E}_{1} - \nu$	18400	17100	17300	cm -1
δ	0.56	0.47	0.48	
$(n_N^+ \rightarrow 4p)$ calcd	63000	57500	48500	cm -1
<i>v</i> <sub>2</sub>	63000-64000	58000 <i>a</i>	49000 <i>a</i>	cm <sup>-1</sup>

<sup>a</sup> Corrected for tentative deconvolution (cf. Figure 4 and Table II).

mean first uv excitation energy of trisilacycloheptanes<sup>26</sup> to  $\nu_{\rm m} \sim 46000 \ {\rm cm^{-1}}$ . Substituting these values into the Rydberg formula yields a term value  $R/(n - \delta)^2 \sim 20000$ cm<sup>-1</sup>, which could correspond to a 4psi Rydberg state.<sup>24</sup> If



so, then under local  $C_{2v}$  symmetry only the two transitions  $\sigma_{SiSiSi} \rightarrow 4p_{Si}$  in the SiSiSi plane would be allowed (trisilacycloheptanes:  $\epsilon_{max} = 6700-8700 \text{ l.}/(\text{mol cm})^{26}$ , fitting nicely into the suggested three-membered ring transition state model. Proof or disproof of assumptions like the ones above is presently attempted by combined PE and uv spectroscopy.29

#### **Experimental Section**

Syntheses. All compounds have been prepared following litera-ture routes: bis(trimethylsilyl)diimine,<sup>30</sup> 1-trimethylsilyl-2-tertbutyldiimine,<sup>31</sup> bis(*tert*-butyl)diimine,<sup>32</sup> phenylazo-*tert*-butyl,<sup>31</sup> phenylazotrimethylsilyl,<sup>31</sup> phenylazotrimethylgermyl.<sup>31</sup> Their purity has been controlled by standard techniques.

Spectra. The PE spectra were recorded using a Perkin-Elmer PS 16. Due to the thermal instability-especially of bis(trimethylsilyl)diimine-the compounds were evaporated directly into the ionization chamber from a bulb cooled by a dry ice bath. Regulating the height of the dry ice bath allowed the maintenance of an approximately constant vapor pressure of the substance. This technique, however, unfortunately did not yield high quality spectra, and therefore ionization energies are only reliable to 0.1-0.2 eV. Uv spectra were recorded in the gas phase on a McPherson 224 equipped with a Hinteregger hydrogen lamp and in solution on a Cary 14. All hydrocarbon solvents have been of spectral purity.

Calculations. The CNDO calculations have been performed at the Hochschul-Rechenzentrum at Frankfurt University, using a modified version<sup>33</sup> of the QCPE program 141. Bond lengths of bis-(trimethylsilyl)diimine have been taken from the structure determination,<sup>13</sup> whereas the bond angles and dihedral angles have been idealized. For the di-tert-butyl and the unsymmetrically substituted compound, standard geometries have been used. The uv transition energies have been calculated by adding a subroutine, where  $K_{ik}$  and  $J_{ik}$  are computed from the CNDO LCAO expansion coefficients and the CNDO atomic electron-electron repulsion integrals  $\gamma_{AB}$ .<sup>23</sup>

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# Reaction of Thiol Anions with Benzene Oxide and Malachite Green

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Abstract: The second-order rate constants for the reactions of a number of thiol anions with benzene oxide (BO) have been determined as have the rate constants and equilibria for the reaction of thiol anions with the acid-base equilibria species of malachite green (MG).  $\beta$  constants for reaction of thiol anion and amines with the nonprotonated species of malachite green are comparable ( $\sim 0.3$ ) to the  $\beta$  constants for the reactions of amines and thiol anions with ethylene oxide (EO) and thiol anions with BO. In addition the rate constants for the reaction of thiol anions with BO and EO are comparable. These results are discussed in terms of attack of RS<sup>-</sup> directly upon BO and EO in an SN2 reaction vs. trapping of intimate ion pairs by RS<sup>-</sup>. As compared with the ability of ethylene oxide to react with nucleophiles possessing basicities as low as that of water, benzene oxide is attacked by only the more polarizable nucleophile as  $N_3^-$ , RS<sup>-</sup>, etc. This is shown to be due not to the greater selectivity of benzene oxide toward soft bases, but to the fact that the spontaneous aromatization rate (i.e., water rate) of benzene oxide exceeds the rate of reaction of ethylene oxide with water by ca. 10<sup>3</sup>. It is predicted that benzene oxides substituted with highly electron-withdrawing groups will exhibit reactivity to a broad range of nucleophiles. This prediction is predicated on the previously determined fact that aromatization of benzene oxide is associated with a  $\rho \simeq -7$ . For the purposes of this study, it was desirable to determine the microscopic  $pK_a$  values for the thiol species under the conditions of the kinetic and thermodynamic studies (30°, solvent water,  $\mu = 1.0$ ). These constants are reported herein and compared with literature values.

As part of the normal metabolic paths of catabolism and detoxification, both biotic and exobiotic aromatic structures are converted to arene oxides. However, these normal metabolic processes may result in the assassination of the subject. Thus, arene oxide moieties formed from exobiotic aromatic hydrocarbons have been implicated as causative agents in mutagenesis, carcinogenesis, and tissue necrosis.<sup>2,3</sup> The pathological effects of arene oxides have been attributed to their acting as alkylating agents.4-7 The enzyme<sup>8</sup> and nonenzyme mediated 1,2-addition of the sulfhydryl group of glutathione to arene oxides appears as a major

means of protection from the cytotoxic effects of these

agents. Thus, both the cytotoxic effects of arene oxides and

the detoxification of these substances involve nucleophilic

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