Infrared Spectra of New Sulfur–Nitrogen Species in Solid Argon

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Abstract: New sulfur–nitrogen species were produced in an argon/nitrogen/sulfur vapor microwave discharge, and the reaction products were trapped in solid argon at 12 K. Nitrogen-15 and sulfur-34 isotopic substitutions, concentration variation, photolysis, and changes upon annealing provided a basis for identification and characterization of the new products. A product absorption at 1209.4 cm⁻¹ is identified as the diatomic SN radical. The strongest product absorption at 1225.2 cm⁻¹, which shows a triplet and a doublet pattern with mixed sulfur and mixed nitrogen isotopes, respectively, is assigned to the antisymmetric stretching vibration of the NS₂ radical; these isotopic data provide a 133 ± 5° measure of the valence angle in NS₂. The less stable NSS isomer absorbs at 1017.1 and 594.6 cm⁻¹ and is converted to NS₂ with near-ultraviolet photolysis. An absorption band at 2040.2 cm⁻¹ shows quartet and doublet patterns with mixed nitrogen and sulfur isotopes, respectively, and is assigned to the nitrogen–nitrogen stretching fundamental of N₂S. Evidence for higher nitrogen sulfides is also presented.

Introduction

Nitrogen oxides have been studied extensively by infrared absorption spectroscopy in the gas phase and solid state. However, the analogous molecular nitrogen sulfides are not well-known. There are, however, sulfur nitrides, which include the polymeric material polythiazyl (SN), and its evaporation products S₂N₂, S₃N₃, S₄N₄, and S₅N₅. This polymeric material has the unusual property that it can be sublimed. An argon matrix infrared study has shown that the S₂N₂ and S₃N₃ molecules evaporate from the polymer. Finally, gas-phase reactions of atomic nitrogen with sulfur compounds have produced solid material containing the above sulfur nitride species.

The reactive SN radical has been widely studied in the gas phase by EPR, microwave, die laser, high-resolution FTIR, electronic absorption, photoelectron, and far-infrared LMR spectroscopies. Its fundamental vibrational band has recently been observed by tunable diode laser at 1204.089 cm⁻¹ for the 1.; ground state in the gas phase. N₂S was first suggested as the carrier for CS₂-N₂ laser action, and its gas-phase N–N stretching frequency was predicted by comparison with N₂-O to be about 300 cm⁻¹ below the 2331-cm⁻¹ vibration of the NS₂ radical; these isotopic data provide a 133 ± 5° measure of the valence angle in NS₂. The less stable NSS isomer absorbs at 1017.1 and 594.6 cm⁻¹ and is converted to NS₂ with near-ultraviolet photolysis. An absorption band at 2040.2 cm⁻¹ shows quartet and doublet patterns with mixed sulfur and mixed nitrogen isotopes, respectively, and is assigned to the nitrogen–nitrogen stretching fundamental of N₂S. Evidence for higher nitrogen sulfides is also presented.

Experimental Section

The vacuum system and chamber for matrix-isolation studies have been described previously. A closed-cycle refrigerator (CTI Cryogenics, Model 22) and an indicator/controller were used to cool and monitor the temperature of the CsI window. FTIR spectra were recorded on a Nicolet 60SX spectrometer at 0.5 cm⁻¹ resolution (KBr beam splitter) and ±0.1 cm⁻¹ frequency accuracy.

Natural sulfur (Electronic Space Products, Inc., recrystallized), enriched sulfur-34 material with 98% 34S, EG&G Mound Applied Technologies, natural isotopic nitrogen gas (Matheson), and 99% nitrogen-15 enriched gas (Cambridge Isotope Laboratories) were used as received. Polymeric sulfur nitride was kindly provided by MacDiarmid. The quartz tube for seeding sulfur vapor into a nitrogen/argon discharge was similar to the one described previously except that the orifice was constricted to 1 mm. The discharged gas stream was condensed at 12 ± 1 K, and FTIR spectra were recorded before and after annealing and photolysis with a medium-pressure mercury arc (Philips, H39KB, 175 W) using long-wavelength pass filters.


In another experiment the sulfur reservoir temperature was increased to 70 °C to supply a 2% nitrogen/argon discharge. An excess of sulfur was evidenced by the observation of weak S\textsubscript{2} and S\textsubscript{3} bands. The relative intensity of S\textsubscript{2}O to SO bands was further enhanced, and the 1225.2 to 1209.4 cm\textsuperscript{-1} absorbance ratio was increased to 10/1. The 1167.4- and 594.6-cm\textsuperscript{-1} bands were enhanced by factors of 10 and 15, respectively. The 2449.0-, 2040.2-, and 1884.9-cm\textsuperscript{-1} bands were also increased. The 773.0- and 882.3-cm\textsuperscript{-1} bands were also produced in very small quantity with a relative intensity of 2/1. New weak absorption bands were observed at 1499.7, 1309.4, 1017.1, and 624.2 cm\textsuperscript{-1}.

Table I. Product Absorptions (cm\textsuperscript{-1}) Observed from the Matrix Reaction between Atomic Sulfur and Nitrogen at Several Sulfur Reservoir Temperatures

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>2449.0</th>
<th>2040.2</th>
<th>1884.9</th>
<th>1499.7</th>
<th>1309.5</th>
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<tr>
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<td>2040.2</td>
<td>1884.9</td>
<td>1499.7</td>
<td>1309.5</td>
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Table II. The Photochemistry and Annealing Behavior of the Product Absorptions (cm\textsuperscript{-1}) Observed from the Matrix Reaction between Atomic Sulfur and Nitrogen

<table>
<thead>
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<th>Band</th>
<th>470 nm</th>
<th>420 nm</th>
<th>340 nm</th>
<th>290 nm</th>
<th>254 nm</th>
<th>20 K</th>
<th>23 °C</th>
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<td>-40</td>
<td>-20</td>
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</table>

a Changes in ± percent of band absorbance. b This band appeared after annealing.
appearance of a weak new band at 893.9 cm\(^{-1}\). The nitrogen concentration was varied from 0.8 to 5%, and 2% was found to be optimum for the maximum yield of sharp new product absorptions.

The spectra obtained from \(^{34}\)S-enriched sulfur with \(^{14}\)N\(_2\) before and after full arc photolysis and annealing at 23 ± 2 K are shown in Figure 2. The changes upon photolysis and annealing are also collected in the Table II. Under similar conditions, several experiments were carried out with \(^{34}\)S-enriched sulfur and \(^{15}\)N-enriched nitrogen. The spectra obtained from the reactions of natural isotopic nitrogen with natural isotopic sulfur, 98% and 50% \(^{34}\)S-enriched sulfur, as well as 99% and 50% \(^{15}\)N-enriched nitrogen with natural isotopic sulfur are shown for the 2100-1400- and 1400-500-cm\(^{-1}\) spectral regions, respectively, in Figures 3 and 4. The isotopic counterparts for all of the observed sulfur-nitrogen species are collected in Table III. Another experiment was done with \(^{14}\)N\(^{15}\)N-enriched gas; the isotopic distribution was the same as with \(^{14}\)N\(_2\) and \(^{15}\)N\(_2\) mixtures, indicating that the nitrogen discharge reaction with sulfur produced atomic nitrogen species.

Oxygen (0.3%) was added to the nitrogen (2%) mixture in one study. Some of the very low intensity bands were enhanced whereas the other bands with relatively higher absorption intensities were decreased. The bands enhanced by the addition of oxygen will be presented in a separate communication.

An early experiment employed a 6-mm open-end discharge tube.\(^{27}\) The overall yield was lower than that obtained from the 1-mm-orifice discharge tube used for the rest of these experiments.

Polymeric Sulfur Nitride. Sulfur nitride (SN), was put in the sample reservoir of the 6-mm open end discharge tube and heated to 130 \(^\circ\)C while 1 mmol/h of argon was allowed to sweep the SN vapor. Most of the sample was deposited within the front unheated part of the tube, and only a small portion reached the cold window. The spectrum was similar to that reported from codeposition of argon with (SN), heated within a Knudsen cell at 160 \(^\circ\)C,\(^{27}\) except that no absorption around 1225 cm\(^{-1}\) was observed. In our experiment S\(_3\)N\(_4\) was observed at 554, 706, and...
Figure 2. Infrared spectra in the 1600–500-cm⁻¹ region for the discharged sulfur vapor/nitrogen/argon stream deposited on a CsI window at 12 ± 1 K: (a) after 6 h of deposition; (b) after 30 min of photolysis with the full arc lamp (254 nm), and (c) after annealing for 15 min at 30 ± 2 K. These spectra were obtained with 98% enriched sulfur-34.

Table III. Product Absorptions (cm⁻¹) Observed from the Matrix Reaction between Atomic Sulfur and Nitrogen

<table>
<thead>
<tr>
<th>²³S</th>
<th>³²S/³⁴S</th>
<th>³⁴S</th>
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<td>1279.5</td>
<td>1272.9</td>
<td>1267.5</td>
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<td>SNSS or NSSS (S-S)</td>
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*Mixed nitrogen isotopic bands are 2009.3 and 2003.9 cm⁻¹. These bands were calculated from ν₁ and ν₁ + ν₂.

948 cm⁻¹ and S₉N₉ showed absorptions at 472, 566, and 788 cm⁻¹ based on comparison to gas-phase spectra. A Tesla coil argon discharge flow was passed over the deposited film of (SN)₉ inside of the tube for 30 min, and the S₉N₉ bands were enhanced. Photolysis of the matrix with 254-nm light caused the S₉N₉ bands to decrease and a new band appeared at 519 cm⁻¹; annealing at 23 ± 2 K produced no new bands and caused the S₉N₉ species to aggregate.

In another experiment, (SN)₉ vapor diluted in argon was subjected to a 40–80-W microwave discharge. With a 130 °C sample reservoir only a weak 1225.2-cm⁻¹ band was observed, and with the temperature increased to 160 °C, S₉ bands were also produced. In the final experiment with (SN)₃, the temperature was kept at 160 °C and the microwave power at 5–10 W. Under these conditions a spectrum similar to the 2% mixture of nitrogen/argon with 70 °C sulfur reservoir was obtained. Similar photolysis and annealing behavior was also observed in this experiment. However, a different intensity distribution was noticed for several bands. The major features of the spectrum compared to that obtained from the mixture of nitrogen and sulfur (70 °C) discharge were the following: (a) a relatively higher yield of 775.0, 882.3, 1309.5, 1253.8, and 624.2-cm⁻¹ bands; (b) the
SN Radical. The absorption band at 1209.4 cm\(^{-1}\) is due to one of the simple primary products in the reaction of nitrogen with dilute sulfur. The 1209.4 cm\(^{-1}\) band showed doublet patterns for both cases of 50:50 mixed \(^{32}\)S\(^{34}\)S and \(^{14}\)N:\(^{15}\)N samples, indicating that this vibration involves only one sulfur atom and one nitrogen atom. The observed sulfur and nitrogen isotopic ratios are 1209.4/1198.6 = 1.0090 and 1209.4/1181.3 = 1.0238, respectively. These values are in excellent agreement with the values of 1.0091 and 1.0240 calculated for a diatomic N=S harmonic oscillator. This absorption is also in excellent agreement with the 1204.089 cm\(^{-1}\) absorption of the diatomic SN in the gas phase.\(^{16,17}\) The SN molecule is similar to PS in that it is not photosensitive and decreases upon annealing to form higher sulfides.\(^{28}\) Although PS sustains a red shift of 3.9 cm\(^{-1}\) in solid argon, SN sustains a 5.3 cm\(^{-1}\) blue shift.

NS\(_2\). The species absorbing at 1225.2 cm\(^{-1}\) was also a simple primary product formed in the reaction of nitrogen with dilute sulfur, and it increased relative to SN as the concentration of sulfur was increased. It was also the major species produced when polymeric sulfur nitride was passed through the 40–80-W microwave discharge. The 1209.4 cm\(^{-1}\) band showed a symmetrical 1:2:1 triplet pattern with a 50:50 mixed \(^{32}\)S:34\(^{15}\)N sample and a doublet pattern with a 50.50 \(^{14}\)N:34\(^{15}\)N-enriched nitrogen, indicating a vibrational motion involving one nitrogen and two equivalent sulfur atoms. This band is assigned to the antisymmetric stretching vibration of S-N-S. The absorption bands at 1884.9 and 2449 cm\(^{-1}\), which were produced in all of the experiments and show the same annealing and photolysis behavior, are assigned to the \(v_1 + v_3\) combination and \(2v_2\) overtone bands, respectively. The symmetric stretching mode, which is expected to be a weak band near 660 cm\(^{-1}\), cannot be observed due to spectral interference from sulfur species in this region. The isotopic shifts for the calculated \(v_1\) band (659.7 cm\(^{-1}\)) are in accord with a symmetric S-N-S stretch.

The valence angle for the C\(_3\) S-N-S molecule can be calculated from pairs of isotopic molecules. As has been discussed for SO\(_2\), central isotopic substitution gives a lower limit and terminal substitution an upper limit to the bond angle owing to the effect of normal cubic anharmonicity.\(^{29}\) The average value with anharmonicity cancellation provides an excellent measure of the valence angle. In the case of O\(_2\), the lower limit 108 ± 1° and upper limit 124 ± 1° averaged to 116 ± 2°, which is in agreement with the microwave value.\(^{30}\) In the case of S\(_2\), the lower limit 114 ± 1° and the upper limit 118 ± 1° averaged to 116 ± 2°, which is in agreement with high-level theoretical calculations.\(^{26}\)

In the present NS\(_2\) case the 32-14-32/32-15-32 isotopic pair gives 140.7° and the 34-14-34/34-15-34 isotopic pair gives 137.7° for upper limits; the upper limit from the 32-14-32/32-15-34 pair is 172.7° and from the 32-15-32/34-15-34 pair is essentially 180° for the valence angle of NS\(_2\). Since we are in fact calculating the cosine of the valence angle from the isotopic frequency ratios, the average of the four cosine values gives 0.892 or an angle of 153°. We prefer the determination 153 ± 5° for the valence angle of NS\(_2\).

The matrix characterization of NS\(_2\) is in excellent agreement with detailed high-level theoretical calculations.\(^{26}\) Depending on the basis set and the level of theory employed, the calculated valence angle ranged from 150 to 155°, which is in excellent agreement with the 153 ± 5° value determined from matrix isotopic frequencies. The single and double configuration interaction calculations, expected to be the most reliable by the authors,\(^{26}\) predict a 151 ± 1° valence angle and frequencies near 730, 300, and 1299 cm\(^{-1}\). Ab initio frequencies are well-known to exceed experimental values, and scale factors near 0.90 are often employed. Scaling the 730-cm\(^{-1}\) calculated value predicts \(v_1\) at 660 cm\(^{-1}\), precisely the value determined from the \(v_1\) and \(v_3\) bands in the matrix spectrum. Scaling the 1299-cm\(^{-1}\) calculated value predicts \(v_1\) at 1169 cm\(^{-1}\), somewhat below the observed 1225-cm\(^{-1}\) value. This discrepancy shows that different scale factors are required for different modes. A scale factor of 0.94 is required to fit the calculated and observed \(v_1\) values for NS\(_2\), which is within the range of acceptable values. Finally, the CASSCF calculations\(^{26}\) predict frequencies that are too low even without scaling; these calculations are clearly inferior to the excellent DZ+P (CISD) calculations for NS\(_2\). Finally, comparison of the overtone (2\(v_2\)) and fundamental (\(v_2\)) shows that this mode

Figure 3. Infrared spectra in the 2100–1400 cm\(^{-1}\) region for the discharged sulfur vapor/nitrogen/argon stream deposited on a CsI window at 12 ± 1 K: (a) natural isotopic sulfur, (b) 98% \(^{34}\)S-enriched sulfur, (c) 50% \(^{34}\)S-enriched sulfur, (d) 99% \(^{15}\)N-enriched nitrogen, and (e) 50% \(^{15}\)N and 50% \(^{14}\)N. (The microwave discharge randomly enriched the isotopic products of the sulfur reaction.)

Discussion

New species will be identified from chemical information and sulfur and nitrogen isotopic data. The simple binary nitrogen sulfides were prepared by reacting atomic nitrogen and sulfur, and as the concentration of sulfur was increased, higher nitrogen sulfides were formed. Secondary reaction products involving the dilute sulfur. The 1209.4-cm\(^{-1}\) band showed doublet patterns for charged sulfur vapor/nitrogen/argon stream deposited absorption bands showed the same photolysis and annealing behavior, are assigned to the \(v_1 + v_3\) combination and \(2v_2\) overtone bands, respectively. The symmetric stretching mode, which is expected to be a weak band near 660 cm\(^{-1}\), cannot be observed due to spectral interference from sulfur species in this region. The isotopic shifts for the calculated \(v_1\) band (659.7 cm\(^{-1}\)) are in accord with a symmetric S-N-S stretch.

The SN molecule is similar to PS in that it is not photosensitive and decreases upon annealing to form higher sulfides.\(^{28}\) Although PS sustains a red shift of 3.9 cm\(^{-1}\) in solid argon, SN sustains a 5.3 cm\(^{-1}\) blue shift.

The absorption band at 1209.4 cm\(^{-1}\) is due to one of the simple primary products in the reaction of nitrogen with dilute sulfur. The 1209.4 cm\(^{-1}\) band showed doublet patterns for both cases of 50:50 mixed \(^{32}\)S:34\(^{15}\)N and \(^{14}\)N:\(^{15}\)N samples, indicating that this vibration involves only one sulfur atom and one nitrogen atom. The observed sulfur and nitrogen isotopic ratios are 1209.4/1198.6 = 1.0090 and 1209.4/1181.3 = 1.0238, respectively. These values are in excellent agreement with the values


Figure 4. Infrared spectra in the 1400-500 cm\(^{-1}\) region for the discharged sulfur vapor/nitrogen/argon stream deposited on a CsI window at 12 ± 1 K: (a) natural isotopic sulfur, (b) 98\% \(^{34}\)S-enriched sulfur, (c) 50\% \(^{34}\)S-enriched sulfur, (d) 99\% \(^{15}\)N-enriched nitrogen, and (e) 50\% \(^{15}\)N\(_2\) and 50\% \(^{14}\)N\(_2\).

is nearly harmonic (2449 cm\(^{-1}\) vs 2 \times 1225.2 = 2450.4 cm\(^{-1}\)), which adds credence to this valence angle determination from isotopic fundamentals.

\(\text{N}_2\text{S}\). The absorption band at 2040.2 cm\(^{-1}\), which shows a quartet at 2040.2, 2009.3, 2003.9, and 1972.3 cm\(^{-1}\) with mixed isotopic nitrogen, is 291 cm\(^{-1}\) below the \(\text{N}_2\) fundamental vibration (2331 cm\(^{-1}\)). This vibration is clearly a \(\text{N-N}\) stretching mode based on the 14-14/15-15 ratio of 2040.2/1972.3 = 1.0344 and agreement with the 1.0350 harmonic diatomic \(\text{N}_2\) ratio and the small \(^{34}\)S shift to 2039.9 cm\(^{-1}\). The isotopic data are indicative of the \(\text{N-N}\) vibration of two inequivalent nitrogen atoms coupled very slightly to sulfur, which lead directly to the NNS assignment for the 2040.2-cm\(^{-1}\) absorption. The present argon matrix observation is in near agreement with the gas-phase value of 2047.59 cm\(^{-1}\) for a new species produced by pyrolysis of 5-phenyl-1,2,3,4-thiatriazole.\(^{(31)}\) The previous matrix value\(^{(32)}\) of 2030 cm\(^{-1}\), which was observed after flash vacuum pyrolysis of the same compound, is shifted by interaction with other species including precursor in that experiment. The lack of observation of the \(\text{N-S}\) stretching vibration in this species is attributed to its low intensity.

The \(\text{N}_2\text{S}\) species was detected in the nitrogen–sulfur experiment with the sulfur reservoir temperature of 70 °C and was enhanced in yield as the sulfur reservoir temperature was increased. This suggests that \(\text{N-N}\) bond cleavage is facilitated through chemical reaction between excited \(\text{N}_2\) and sulfur. Finally, the \(\text{N}_2\text{S}\) species was made here from the elements instead of a large cyclic organic precursor, which further supports its identification as a simple molecule.

\(\text{NSS}\). The absorption band at 594.6 cm\(^{-1}\) was not detected in the discharge reaction of nitrogen plus sulfur with the sulfur reservoir at 25 °C; however, it was detected when the sulfur reservoir temperature was increased to 50 °C. Under this condition the total amount of sulfur, especially the ratio of diatomic sulfur to atomic sulfur, increased as was evidenced by the absolute and relative intensity of \(\text{S}_2\text{O}/\text{SO}\) bands, which changed from 1:2 to more than 2:1; however, no \(\text{S}_3\) or \(\text{S}_4\) was found. As the sulfur reservoir temperature was further increased to 70 and 115 °C, the relative intensity of \(\text{S}_2\text{O}\) to \(\text{SO}\) as well as the 594.6 cm\(^{-1}\) band to \(\text{SN}\) (1209.4 cm\(^{-1}\)) increased. A 1:1:1:1 quartet pattern with the inner bands very close to each other was obtained with a 50:50 mixed isotopic sulfur. Mixed isotopic nitrogen produced a doublet pattern with only 1.4-cm\(^{-1}\) spacing. This absorption band is 121.4 cm\(^{-1}\) below the sulfur–sulfur vibration frequency in diatomic \(\text{S}_2\).\(^{(32)}\) The sulfur isotopic ratio of 594.6/578.1 = 1.0285 is very close to the sulfur isotopic value of 672.6/653.7 = 1.0289 in \(\text{S}_2\text{O}\), but less than that for a pure sulfur–sulfur vibration (1.0304), which suggests an S–S species bonded to another atom. Furthermore, the close inner bands in the mixed isotopic sulfur and the small shift with nitrogen-15 (594.6–593.2 = 1.4 cm\(^{-1}\)) indicate that this S–S vibrational mode is only slightly perturbed by the other bonded species. A similar isotopic oxygen shift (672.7–671.0 =


1.7 cm⁻¹) was also observed for the S–S stretching mode of S₂O. These isotopic characteristics of the 594.6-cm⁻¹ band are in parallel with those of SSO. Similarly, the 594.6-cm⁻¹ band is proposed to be the sulfur–sulfur vibration mode in the NSS species.²²

The 1017.1-cm⁻¹ band exhibits a constant absorbance relative to the 594.6-cm⁻¹ band, namely 1/9, in the present experiments with different reagent concentrations, on photolysis, and upon annealing. The sulfur isotopic ratio 1017.1/1008.1 = 1.0090 and the nitrogen isotopic ratio 1017.1/993.1 = 1.0242 are near diatomic SN values, which identifies this band as an S–S vibration. Mixed isotopic experiments reveal doublets for the vibration of single S and N atoms. The 1017.1-cm⁻¹ band is assigned to the N–S stretching mode in the NSS molecule. The asymmetric SSS isomer is predicted to be 19 kcal/mol higher in energy than the CS₂ isomer by the same high-level theoretical calculations.²⁶ The strongest fundamental absorption is predicted for the S–S stretching mode at 665 cm⁻¹. A reasonable scale factor 0.895 takes the calculated 665-cm⁻¹ value into the observed 595-cm⁻¹ value. The calculated S–S bond length once formed in the matrix NSS is stable until energized. Irradiation with near-ultraviolet light activates the NSS isomer and shows a constant absorption intensity ratio of 3/1 in all of the produced only when the concentration of sulfur was increased to allows isomerization to the more stable NS₂ form (Figure 3).

The observed 595-cm⁻¹ value. The calculated S–S bond length of 1.920 Å is longer than the S value of 1.89 Å which has an appropriately higher frequency (720 cm⁻¹).²¹,²² The N–S stretching fundamental is predicted to be half as intense and at 1072 cm⁻¹. A scale factor of 0.949 is required to relate the calculated value to the observed value. Again different scale factors are required for different modes of the same molecule.

The NSS species is probably formed from the reaction of N plus S₂ or SN plus S which also gives SNS. Although NSS is less stable than NS₂,

\[
N + S₂ \rightarrow N-S-S
\]

SN + S \rightarrow N-S-S, S-N-S

once formed in the matrix NSS is stable until energized. Irradiation with near-ultraviolet light activates the NSS isomer and allows isomerization to the more stable NS₂ form (Figure 3).

NSS \rightarrow 400-254 \text{ nm} \rightarrow NS₂

This photoisomerization process is apparently not reversible. Since S₂ absorbs in the near-ultraviolet region, it is reasonable to expect NSS to absorb in the same region.

NSSS. The absorption bands at 1309.5 and 624.2 cm⁻¹ were produced only when the concentration of sulfur was increased to the point that excess sulfur was observed as S₂. The above bands show a constant absorption intensity ratio of 3/1 in all of the experiments. They decreased upon photolysis with 420-nm radiation and gives perturbed SN at 1253.7 cm⁻¹ and another perturbed S₂, which was not observed presumably due to low intensity or masking by S₂ and SNS absorptions. The original NS–S₂ species was reformed upon annealing.

This species is produced in the presence of excess sulfur, manifested as S₂, suggesting that it is probably formed from the reaction of S with nitrogen in the gas phase or on the matrix surface. However, reaction of SN with S₂ is apparently the most probable mechanism within the matrix.

\[
N + S₂ \rightarrow NSSS \text{ (gas phase or on the matrix surface)}
\]

SN + S₂ \rightarrow NSSS \text{ (within matrix)}

The latter pathway is in agreement with the annealing behavior subsequent to the photolysis where adjacent SN and S₂ can react with each other in contrast to the small growth of this species upon direct annealing where only a small amount of perturbed SN with S₂ is present.

SNNS₂. The absorption band at 1499.7 cm⁻¹ was produced as the sulfur reservoir temperature increased to 70 °C and higher. However, annealing decreased this band, indicating a more stable higher sulfide absorbs elsewhere. The fact that the 1499.7-cm⁻¹ band was observed in the presence of a relatively high yield of the primary reaction products SN and NS₂ suggests a secondary reaction product of the primary species. The sulfur and nitrogen isotopic ratios for the upper band are very close to those observed for the antisymmetric stretch of NS₂. We assign the 1499.7-cm⁻¹ absorption band to the strong antisymmetric NS₂ stretching vibration in the SN–NS₂ molecule; the weak N–S stretching mode was not observed. This assignment requires a relatively weak N–N bond. As a consequence the vibrational motion of one fundamental group is not coupled with the vibrational motion of the other group, and no nitrogen isotopic mixing occurs; this is the case for the analogous ON–NO₂ molecule.³⁵

It is interesting to compare the fundamentals in SN–NS₂ and ON–NO₂ with their constituent functional group fundamentals. The major difference here is the electron-withdrawing effect of the –NO₂ group versus the probable electron-donating effect of the –NS₂ group owing to the substantially lower electronegativity of S as compared to O. The frequency shifts are relatively small in ON–NO₂ from the subgroup fundamentals; however, in SN–NS₂, the –NS₂ group fundamental increases 274 cm⁻¹. This denotes antibonding electron density displacement from NS₂ to NS₃ with concomitant displacements in group fundamentals. This hypothesis is supported by the higher calculated electron affinity for NS as compared to NO₃.

\[
\text{NS–SN, the 1167.5-cm}^{-1} \text{ band was formed in all experiments where SN was produced, but this band also decreased on sample annealing. The intensity relative to SN increased from 1/2 with the sulfur reservoir temperature at 50 °C to 1/1 with this temperature at 70 °C. The sulfur and nitrogen isotopic ratios for this band are in very close agreement with diatomic SN with the sulfur ratio slightly higher and the nitrogen ratio slightly lower than those of SN.}
\]

A missing species is (NS₃). The cyclic S₃N₂ molecule evaporation from the solid is clearly not produced from the elements in a gaseous discharge as S₃N₂ was not observed in the present experiments. Although S₃N₂ is a stable vapor species, it must be formed by evaporation or reaction of the solid S₃N₄ compounds.³⁴ The 1167.5-cm⁻¹ band is tentatively assigned to NS–SN bonded through sulfur in contrast to (NO₂), which is weakly bonded through nitrogen. Here the increased importance of sulfur–sulfur bonding is noteworthy.

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NS$_2$. The weak 893.9-cm$^{-1}$ band was formed only after annealing the matrix and shows doublet and triplet patterns with mixed isotopic nitrogen and sulfur, respectively. The sulfur and nitrogen isotopic ratios are in very close agreement with those found for the NS$_2$ molecule with the sulfur ratio slightly higher and the nitrogen ratio slightly lower than those of NS$_2$. This 893.9-cm$^{-1}$ band also decreased upon subsequent photolysis with the 254-nm light. We assign this band to the antisymmetric stretching vibration of NS$_2$. The bond angle lower limit for this C$_S$ species calculated from the 32-14-32/32-15-32 isotopic pair is 113.2° and that from the 34-14-34/34-15-34 isotopic pair is 112.9°. The upper limit from 32-14-32/34-14-34 is 121.2° and that from 32-15-32/34-15-34 is 122.6°. The average bond angle calculated from the isotopic ratios for NS$_2$ is 117 ± 3°. As expected this angle is substantially smaller than the 153 ± 5° value determined for NS$_2$. Similar relationships have been found for the anion/neutral pairs NO$_2$/NO$_{3}^-$, PS$_2$/PS$_{3}^-$, PO$_4$/PO$_{3}^-$, and O$_3$/O$_{3}^-$.30, 40

Since the 1-mm-orifice discharge tube was used in these experiments, charged species are expected to play a subordinate role compared to neutral species; however, the NS$_2$ absorption at 1225.2 cm$^{-1}$ was very strong, and NS$_2$ is the most likely anion to give a strong infrared signal in these experiments. Diffusion and reaction of SN and S is the most likely source of NS$_2$ in these experiments.

NS$_2S$. The relative intensity of the absorption bands at 773.0 and 882.3 cm$^{-1}$ remains constant in all of the experiments, and both decrease on 340- and 290-nm photolysis and increase upon 254-nm photolysis and increase on annealing with the same relative intensity. This behavior suggests that these two bands belong to the same species formed from reaction of two other primary species or rearrangement of another isomer. The fact that these bands were detected in the 50 °C sulfur experiment after annealing supports formation of these bands from the primary species SN or NS$_2$. The sulfur triplet isotopic pattern for both bands also suggests the presence of two nearly equivalent sulfur atoms in both of these vibrational motions. The sulfur and nitrogen isotopic ratios for the 882.3-cm$^{-1}$ band are intermediate between the ratios for NS$_2$ and SN, which is indicative of a valence angle less than the value calculated above the NS$_2$ if this is an antisymmetric N-S$_2$ stretching mode. On the other hand, the 773.0-cm$^{-1}$ band exhibits the opposite trend in isotopic ratio which is in accord with a symmetric N-S$_2$ stretching mode.

The vibrational data indicate antisymmetric and symmetric vibrations of NS$_2$ group(s) with equivalent sulfur atoms, but the region of these absorptions is characteristic of N-S single bonds (i.e. S$_2$N$_2$ and S$_2$N$_2$) instead of an isolated -NS$_2$ group (NS$_2$ radical absorbs at 1225 cm$^{-1}$). The above observations suggest a NS$_2$ radical dimer species (NS$_2$)$_2$ in a six-membered ring structure with two S-S single bonds and no N-N bond. This is in marked contrast with the structure of N$_2$O$_4$, which has a weak N-N bond, but it is similar to the structure of S$_2$N$_2$, a minor compound in the thermal decomposition of (SN)$_x$.8 The previously characterized S$_2$N$_2$ molecule forms a puckered six-membered ring with S atoms in the 1 and 3 positions and has strong infrared absorptions at 1034 and 628 cm$^{-1}$. The present suggested NS$_2$ radical dimer ring NS$_2$M$_2$ molecule has N atoms in the 1 and 4 positions and absorptions at 882.3 and 773.0 cm$^{-1}$. Since S-S bonds are known to be stronger than O-O bonds, in the above suggested NS$_2$M$_2$, the NS$_2$ subgroups are united by two stronger S-S bonds instead of one weak N-N bond. Finally, this NS$_2$ species appears to be the most stable final product of the reaction of excess sulfur and nitrogen atoms. It would be interesting to compare theoretical calculations of 1,3- and 1,4-tetrasulfur dinitride rings.

**Reaction Mechanisms.** Nitrogen discharge has been widely studied both for fundamental reasons and practical applications, and the role of N atoms and N$_2$.4 metastable molecules in "active nitrogen" has been considered.45-49 It is generally believed that excitation of N$_2$ to metastable state N$_2^*$ is easier than dissociation to N atoms and that dissociation follows electron impact of N$_2^*$. Accordingly, the concentration of species present in the microwave discharge decreases in the order Ar > N$_2$ > N$_2^*$ > N. Flowing microwave discharge systems are known to contain approximately 10$^{12}$ N atoms/cm$^{-3}$.55 Under the present pressure conditions, approximately 10$^5$ atoms/cm$^3$ are present, but only 2% of these are N$_2$. If the same N atom concentration is attained, then 1 in 20 N$_2$ molecules is dissociated. It is, however, known that N atoms are effective in relaxing N$_2^*$ which provides a mechanism for isotopic scrambling in the discharge.55 In the present $^{14}$N$_2$+$^3$N$_2$ experiments, four isotopic N$_2$O$_4$ molecules were observed as a scrambled nitrogen isotopic quartet at 2218.0, 2199.6, 2177.5, and 2149.4 cm$^{-1}$ in solid argon.50 Since sulfur atoms and S$_2$ are also produced in the sulfur-doped discharge,27 the primary reactions to give SN in this system are (1a) and (1b). Since N$_2$ produces atomic N in the discharge, reaction 1a probably provides more SN than reaction 1b.

$$\text{N}_2^* + S \rightarrow \text{NS} + \text{N}$$
$$\text{N} + \text{S}_2 \rightarrow \text{NS} + \text{S}$$

The addition of more sulfur atoms on the matrix surface during condensation, reaction 2, accounts for the formation of NS$_2$, which

$$\text{SN} + \text{S} \rightarrow \text{NS}_2$$

may not survive the discharge. Likewise the addition of nitrogen atoms to SN on the matrix surface, reaction 3, is the preferred

$$\text{SN} + \text{N} \rightarrow \text{N}_2\text{S}$$

mechanism for the formation of N$_2$S. It is, however, possible that reaction 1a produces N$_2$S in the discharge, and a fraction of this may be collisionally relaxed and survive to be trapped in the matrix. Finally, larger species result from combinations of the above primary products during matrix condensation.

**Conclusions.** The argon/nitrogen/sulfur discharge is an effective method for producing nitrogen sulfide species for isolation in solid argon. The parallels with nitrogen oxide chemistry are of most interest; the diatomic and triatomic oxides and sulfides are common, NO, N$_2$O, NO$_2$ and SN, N$_2$S, and NS$_2$, where the major difference is the larger bond angle for NS$_2$ (153 ± 5°) as compared to NO$_2$ (113°). For the larger sulfur-containing molecules, the increased importance of catenation is apparent with sulfur. The less stable NSS isomer is also trapped and photochemically isomerized to NS$_2$, and the addition of another S atom leads to the formation of a chain-like SN-SS species rather than the symmetrical NO$_3$ racemate species. Only one higher sulfide with a -NS$_2$ terminal group was observed, and this is suggested to characterize SN-NS$_2$, which is analogous to ON-NO$_2$. Upon the addition of more sulfur, S-S bonding determines the structure of the proposed NS$_2$ radical dimer (NS$_2$)$_2$, and the absorptions suggest a six-membered ring with two sulfur-sulfur bonds and N atoms in the 1 and 4 positions.

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This differs slightly from the structure of the $\text{S}_3\text{N}_2$ molecule produced from the decomposition of (SN)$_4$. The matrix infrared characterization of larger nitrogen sulfides invites more electronic structure calculations to test these bonding models.

Finally, the gas-phase reaction of N and S atoms at high dilution produced simple nitrogen sulfide species SN, N$_2$S, and NS$_2$ and not the larger sulfide nitride species $\text{S}_3\text{N}_2$ and $\text{S}_4\text{N}_4$ formed on evaporation of solid polymeric sulfur nitride.

An Interpretation of Organometallic Bond Dissociation Energies

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Abstract: Organometallic bond dissociation energies (BDE) are incorporated into the E and C model with excellent results. Since the data are consistent with gas-phase systems, the fit supports claims that the enthalpies are relatively free of solvation contributions. The organometallic catomers (fragments forming the positive end of the dipole) include (CO)$_5$Mn$^+$, [111'-C$_5$H$_5$]$_2$Zr$^-$, and [1111-C$_5$H$_5$]$_2$M$^+$, Ru$^{5+}$, [C$_6$H$_5$]$_2$P$_2$CH$_2$P(C$_6$H$_5$)$_2$CH$_2$P(C$_6$H$_5$)$_2$M$^{5+}$, [C$_6$H$_5$]$_2$Pt$^{5+}$, [C$_6$H$_5$]$_2$Zr$^-$, and [1111-C$_5$H$_5$]$_2$M$^+$-Sn$^-$. The animers (fragments forming the negative end of the dipole) include organics, halogens, and inorganics. In addition to predicting and interpreting enthalpies, the resulting parameters for the radicals provide reactivity scales that can be used to interpret reactivity and physicochemical properties. Significant chemical insight results from the fit of the data to the ECT model. The parameters are used to reinterpret $^{19}$F chemical shifts in (F-Ph)Pt(P$_2$R$_3$)$_{-}X$, in terms of a $\pi$-only Pt-X bond. The earlier interpretation is one of the classic examples used to support metal-ligand a-bonding interactions. Predicted bond energies are analyzed to indicate the metal properties that facilitate the CO insertion reaction. The analysis provides insights into the relative importance of electrostatic and covalent interactions. Further, the analysis shows that simply plotting the data of $\text{L}_2\text{M}-\text{X}$ versus $\text{H}-\text{X}$ gives an incomplete and potentially incorrect interpretation of the data. Another outcome of the analysis is that certain data points are brought into question as well as suggestions for key experiments needed to bring better insight into several systems.

Introduction

Organometallic chemists have had considerable success in the study of model reactions whose mechanisms provide insights regarding important fundamental steps in homogeneous and heterogeneous catalytic processes. As with the catalytic system, minor perturbations, e.g., change to another metal or ligand, often lead to dramatic changes in reactivity. Interest in obtaining a more fundamental understanding of this chemistry has led to a desire to understand the bonding energetics. Such understanding is essential to improve and fine tune catalysts. 

Measuring thermodynamic bond strengths or bond dissociation energies (BDE) for these systems is time consuming. It behooves us to extract the maximum amount of information and insight present in the data. An important component of understanding the data is being able to predict bond energies for new systems, thus eliminating the need for additional tedious measurements. Most attempts to predict and interpret bond dissociation energies involve comparing the measured values for organometallic $\text{L}_2\text{M}-\text{X}$ compounds with those for $\text{H}-\text{X}$, where $\text{X}$ represents the negative end of the bond dipole and can be either an organic or inorganic group. This approach is of limited and uncertain utility because the relative importance of covalent and electrostatic bonding contributions usually varies in the $\text{L}_2\text{M}-\text{X}$ and $\text{H}-\text{X}$ compounds being compared. The complications introduced into such correlations from variation in the covalency have been discussed in detail for neutral donor-acceptor adducts and for gas-phase ion-molecule correlations with proton affinities.

In most correlations, systems that deviate provide the most interesting cases. For example, deviations from a linear plot of the dissociation energies of $\text{L}_2\text{M}-\text{X}$ versus $\text{H}-\text{X}$ are often attributed to $\pi$-back-bond stabilization or steric effects. These conclusions are open to question because deviations from these plots can arise from variation in the relative importance of covalent and electrostatic contributions to the bonding in the two systems plotted. Another approach to explaining deviations in plots of $\text{H}-\text{X}$ bond energies versus $\text{L}_2\text{M}-\text{X}$ is to draw a second line for a new class of $\text{X}$ groups. This is a sign that the covalent contribution to the bonding in $\text{L}_2\text{M}$ is different from that in $\text{H}$. When $\text{H}-\text{X}$ bond energies are plotted versus $\text{L}_2\text{M}-\text{X}$ bond energies, linear plots usually result when $\text{X}$ is limited to a series of halides or to a series of alkyl compounds. When such plots are found, the conclusion that $\text{L}_2\text{M}$ and $\text{H}$ are similar is incorrect. For limited data sets, straight lines can be obtained even if $\text{L}_2\text{M}$ and $\text{H}$ are very different. In general, a plot of bond energies for two different systems provides little insight about the fragments plotted, and when deviations from the plot occur for

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Registry No. S, 7704-34-9; N$_2$, 7727-37-9; SN, 12033-56-6; S$_2$S$_4$, 12033-57-7; N$_2$S, 56400-02-3; N$_2$, 137122-34-0; N$_2$SS, 88574-94-1; SN$_2$, 137122-35-1; NS-SN, 137122-36-2; $\text{N}_2$, 91054-03-4; N$_2$S, 79796-33-1; $^{15}$N, 14390-96-6; $^{18}$N, 13981-57-2; $^{14}$S, 13965-97-4; Ar, 7440-37-1.


(2) Drago, R. S. Inorg. Chem. 1990, 29, 1379. Equation 2 becomes a linear equation for acids with the same $C_1/E_3$ ratio $-\Delta H/E_3 = E_1/E_3 + C_1/C_3 E_3 = E_1 + kE_3$. Thus $-\Delta H = PB_{12}$ where $PB_1$ is a single parameter base scale equal to $E_1$ and $k = C_1/E_3$. If the enthalpies of two acids are linear with the base parameter, they are linear to each other.