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Predicted thermochemistry and unimolecular kinetics of nitrous sulfide

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The geometry of N2S was obtained at the CCSD(T)/aug-cc-pV(T+d)Z level of theory and energies with coupled-cluster single double triple (CCSD(T)) and basis sets up to aug-cc-pV(6+d)Z. After correction for anharmonic zero-point energy, core-valence correlation, correlation up to CCSDT(Q) and relativistic effects, D0 for the N–S bond is estimated as 71.9 kJ mol$^{-1}$, and the corresponding thermochemistry for N2S is $\Delta H_0^\ddagger = 205.4$ kJ mol$^{-1}$ and $\Delta H_{298}^\ddagger = 202.6$ kJ mol$^{-1}$ with an uncertainty of $\pm 2.5$ kJ mol$^{-1}$. Using CCSD(T)/aug-cc-pV(T+d) theory the minimum energy crossing point between singlet and triplet potential energy curves is found at $r(N-N) \approx 1.105$ Å and $r(N-S) \approx 2.232$ Å, with an energy 72 kJ mol$^{-1}$ above N2 + S(3P). Application of Troe’s unimolecular formalism yields the low-pressure-limiting rate constant for dissociation of N2S $k_0 = 7.6 \times 10^{-10}$ exp$\left(\frac{-126 \text{ kJ mol}^{-1}/R/T}{\text{cm}^3 \text{ molecule}^{-1} \text{s}^{-1}}\right)$ over 700–2000 K. The estimated uncertainty is a factor of 4 arising from unknown parameters for energy transfer between N2S and Ar or N2 bath gas. The thermochemistry and kinetics were included in a mechanism for CO/H2/H2S oxidation and the conclusion is that little NO is produced via subsequent chemistry of NNS. © 2011 American Institute of Physics. [doi:10.1063/1.3628521]

I. INTRODUCTION

The chemistry of nitrogen oxide formation in flames has been studied extensively\(^1\) and the major mechanisms are fairly well established. It is generally recognized that formation of NO\(_x\) may arise from fixation of N\(_2\) in the combustion air or oxidation of organically bound nitrogen in the fuel. Homogeneous mechanisms for fixation of N\(_2\) involve the attack of reactive radicals (O, CH, H) on the triple bond in molecular nitrogen. These reactions form either NO or a reactive nitrogen intermediate (N\(_2\)O, NCN, and NNH) that may subsequently be oxidized to NO. Significant influences of small amounts of sulfur on the formation of nitrogen oxides have been observed at elevated temperatures, but the detailed mechanism is unknown.\(^2\) Here we consider possible roles for nitrous sulfide, N\(_2\)S, the sulfur analog of N\(_2\)O, in order to evaluate the possibility of nitrogen fixation via reaction with atomic sulfur in flames.

The N\(_2\)S molecule has been observed via infrared spectroscopy in the gas-phase and in matrix isolation.\(^3\) It has been the object of several computational studies,\(^1\)\(^4\)\(^6\) with the main goal of interpreting the spectra, and which confirm linear NNS as the most stable structure. Here we present an \textit{ab initio} analysis of the thermochemistry of the N\(_2\)S molecule, and estimates of its unimolecular kinetics at elevated temperatures. Our aim is to check for the potential involvement of this species in the coupling of sulfur and nitrogen chemistry in flames and other high temperature systems. Analogous to the thermal decomposition of N\(_2\)O,\(^7\)\(^8\) the dissociation

\[ N_2S \rightarrow N_2 + S \]  

is spin-forbidden, and we expect the kinetics of reaction (1) to be similarly controlled by crossing between the singlet potential energy curve for the reactant and the triplet potential energy curve for the products.\(^9\) In the case of N\(_2\)O the triplet interaction is repulsive so intersystem crossing occurs at an energy significantly above the dissociation energy of N\(_2\)O to ground state products.\(^10\) The 0 K bond dissociation enthalpy of N\(_2\)O is 161 kJ mol$^{-1}$ while the activation energy for its dissociation is some 100 kJ mol$^{-1}$ higher at the high-pressure limit.\(^5\) Several theoretical studies have focused on the spin-forbidden dissociation of N\(_2\)O (see Refs. 10–12 and references therein).

Here we map out the lowest energy singlet and triplet potential energy surfaces for N\(_2\)S and use the results to address the thermochemistry of this molecule, and to make predictions of its unimolecular reaction kinetics and its potential role in combustion. Energy transfer parameters for N\(_2\)S are estimated by analogy to the valence isoelectronic species N\(_2\)O and carbonyl sulfide OCS, for which dissociation measurements are available.

II. METHODOLOGY

Initial analysis of linear N\(_2\)S in the singlet $^1 \Sigma^+$ and triplet $^3 \Pi$ states was carried out using coupled-cluster single double triple (CCSD(T)) theory\(^13\) and the aug-cc-pV(T+d)Z basis set which includes “tight d” functions on sulfur\(^14\) (and not nitrogen, for which this basis is equal to aug-cc-pVTZ (Refs. 15 and 16)). For the singlet species, spin-restricted Hartree-Fock

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TABLE I. Ab initio energies for the N₂S system (all quantities in a.u. unless otherwise noted, 1 a.u. ∼ 2625.5 kJ mol⁻¹).

<table>
<thead>
<tr>
<th>Species</th>
<th>AV(T + d)Z</th>
<th>AV(Q + d)Z</th>
<th>AV(S + d)Z</th>
<th>AV(6 + d)Z</th>
<th>ZPE²</th>
<th>Scalar relativistic²</th>
<th>Core valence⁴</th>
<th>CCSD(T)/CBS(T)⁴</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂</td>
<td>−109.38085</td>
<td>−109.40721</td>
<td>−109.41546</td>
<td>−109.41833</td>
<td>1176</td>
<td>−0.05829</td>
<td>−0.12119</td>
<td>−0.00155</td>
</tr>
<tr>
<td>S</td>
<td>−397.65722</td>
<td>−397.66862</td>
<td>−397.67210</td>
<td>−397.67356</td>
<td>0</td>
<td>−1.07746</td>
<td>−0.42866</td>
<td>−0.00088</td>
</tr>
<tr>
<td>N₂S</td>
<td>−507.06330</td>
<td>−507.10443</td>
<td>−507.11725</td>
<td>−507.12200</td>
<td>1884</td>
<td>−1.13488</td>
<td>−0.55051</td>
<td>−0.00360</td>
</tr>
<tr>
<td>ΔE⁵</td>
<td>66.26</td>
<td>75.09</td>
<td>77.96</td>
<td>79.07</td>
<td>−8.47</td>
<td>−2.30</td>
<td>1.75</td>
<td>3.10</td>
</tr>
</tbody>
</table>

²In cm⁻¹ (see text).
³Sum of mass-velocity and Darwin terms at CISD/cc-pwCVTZ level.
⁴CCSD(T)/cc-pwCVTZ and CCSD(T)/cc-pwCVQZ data were extrapolated to the CBS limit via Eq. (2), and the difference taken between including a frozen core or using all the electrons in the correlation calculations.
⁵Calculated with the cc-pVTZ basis set.
⁶Contribution to energy change for reaction (1) in kJ mol⁻¹.

and CCSD(T) was employed, and spin-unrestricted versions were used for the triplet species. Spin contamination was negligible, with $S(2S^1)$ below 2.03 for the triplet states, which compares well with the ideal value of 2. The geometries of singlet N₂S and N₂ were optimized. Then $r(N–N)$ in the linear N₂S configuration was optimized as a function of various fixed N–S separations, in both singlet and triplet states, to explore the singlet and triplet potential energy curves. These computations were carried out with the GAUSSIAN 03 program suite.¹⁷

The MOLPRO 2006 program¹⁸ was used to derive the N–S bond dissociation enthalpy in N₂S. At the CCSD(T)/aug-cc-pV(T + d)Z geometries, energies of singlet N₂S, singlet N₂, and triplet S were obtained with a sequence of aug-cc-pV(n + d)Z basis sets with n = 3–6 and CCSD(T) theory (RCCSD(T) for the closed shell species and UCCSD(T) (Ref. 19) for atomic sulfur, all with spin-restricted Hartree-Fock wave functions). Core-valence correlation was assessed with CCSD(T)/cc-pwCVTZ theory²⁰ as the difference between the energy with all electrons correlated and with the core electrons frozen. Convergence of this contribution was investigated by carrying out CCSD(T)/cc-pwCVQZ calculations as well. Scalar relativistic effects were incorporated at the CISD/cc-pwCVTZ level as the sum of the mass-velocity and Darwin terms,²¹ and vector relativistic effects were included via the experimental spin-orbit splitting of the 3P state of atomic sulfur²² (−2.35 kJ mol⁻¹). The MRCC program by Kállay²³ interfaced to the CFour code²⁴ was employed to test the effect of higher level electron correlation by means of CCSD(T) and CCSDT(Q) calculations with the cc-pVTZ basis set.¹⁵,²⁵

III. RESULTS AND DISCUSSION

A. Geometry and thermochemistry

Our $r_e$ for N₂ at the CCSD(T)/aug-cc-pVTZ level of theory is 1.104 Å (1 Å = 10⁻¹⁰ m), which is 0.006 Å greater than the experimental value.²⁶ Combined with the force constant this bond extension would lead to an energy $\sim 0.3$ kJ mol⁻¹ too high relative to the minimum. For N₂S, our CCSD(T)/aug-cc-pV(T + d) computation for $r_e$ (N–N) of 1.132 Å is 0.001 Å above the best estimate of Pak et al.,⁶ which would induce a negligible energy error, less than 0.02 kJ mol⁻¹. It has been noted previously that $r_e$(N–S) tends to be overestimated,⁵,⁶ and indeed our $r_e$(N–S) of 1.593 Å is somewhat too high, and lies 0.015 Å above the best estimate of Pak et al.⁶ Coupled with the N–S stretching force constant this will lead to an energy error of $+0.3$ kJ mol⁻¹, and therefore there is fortuitous cancellation of the contributions of geometry errors to the N₂–S bond dissociation enthalpy.

Table I summarizes the ab initio data for N₂S, N₂, and S, and the equilibrium N₂–S dissociation energy $D_e$ (shown as ΔE in the first four columns) which does not include zero-point vibrational energy. It may be seen that application of large basis sets has almost but not quite converged $D_e$. Several functions have been proposed for extrapolation of individual energies to the infinite or complete basis set (CBS) limit, as a function of an index n which varied from 3 in the aug-cc-pV(T + d)Z basis set through 6 in the aug-cc-pV(6 + d)Z basis. We have employed the two-point relationship of Helgaker et al.,²⁷

$$E_n = E_{CBS} + \frac{B}{n^3}$$

the exponential relation of Feller²⁸ fitted through three points

$$E_n = E_{CBS} + B \exp(-Cn)$$

and the function of Martin²⁹ fitted through all four points

$$E_n = E_{CBS} + \frac{B}{(n+0.5)^4} + \frac{C}{(n+0.5)^6}$$

to extrapolate the total energies (Hartree-Fock plus correlation). The CBS results are shown in Table II. The possible extrapolations involving the largest basis set (n = 6) are Eq. (2) applied to n = 5 and 6, Eq. (3) with n = 4–6, and Eq. (4) (which we fitted to n = 3–6). They lead to $D_e$ = 80.58, 79.62, and 80.23 kJ mol⁻¹, respectively. The mean and standard deviation of these three values are 80.1 and 0.4 kJ mol⁻¹, respectively. We note that the result of Eq. (4) lies in the middle. The two-point scheme with successively larger basis sets approaches this value from above while the two exponential scheme approaches this value from below (see Table II). These trends suggest that the true infinite basis set limit has been bracketed. Error limits of twice the standard deviation should conservatively encompass any uncertainty in the extrapolation. We can use this $D_e$ result to comment on extrapolations relying on smaller basis sets which of course are
dramatically more economical in terms of computer resources. The two-point 3–4 and 4–5 extrapolations give $D_e = 81.5$ and 81.0 kJ mol$^{-1}$, and the exponential extrapolation based on $n = 3–5$ yields $D_e = 79.3$ kJ mol$^{-1}$. These results indicate that for accuracy to within 2 kJ mol$^{-1}$, calculations with up to CCSD(T)/aug-cc-pV(Q+d)Z and a two-point extrapolation would be adequate for the present system.

The core-valence correction to reaction (1) was found to be 1.96 kJ mol$^{-1}$ with the cc-pwCVTZ basis set, and 1.84 kJ mol$^{-1}$ with the cc-pwCVQZ basis set, so it is almost converged with respect to basis set size. The data in Table I were obtained by extrapolating the frozen-core and all-electron results with each basis set to the CBS limit by means of the two-point relation (2), and yield an estimated core-valence correction of 1.75 kJ mol$^{-1}$ at the CBS limit. The equilibrium $D_e$ is then corrected (see Table I) for the core-valence and scalar relativistic effects (which almost cancel), the spin-orbit correction of atomic S($^3P$), and the effect of core-valence and scalar relativistic effects (which almost cancel) to obtain $D_e = 79.3$ kJ mol$^{-1}$. For comparison, this is less than half $D_e$ for N$_2$–O. The tabulated heat of formation of atomic S is 71.9 kJ mol$^{-1}$, and 81.5 and 81.0 kJ mol$^{-1}$ for N$_2$–S, and H$_298$. The tabulated thermal corrections for S and N$_2$, and H$_298$–H$_0$ = 10.26 kJ mol$^{-1}$ computed here for N$_2$–S from the fundamental frequencies recommended by Pak et al.,$^6$ to obtain $\Delta H_0$ = 274.9 kJ mol$^{-1}$ at 0 K, but Nagy et al.$^32$ recently argued for revision to 277.3 kJ mol$^{-1}$. We use the latter value plus tabulated thermal corrections for S and N$_2$, and H$_298$–H$_0$ = 10.26 kJ mol$^{-1}$ computed here for N$_2$–S from the fundamental frequencies recommended by Pak et al.,$^6$ to obtain $\Delta H_0$ = 274.9 kJ mol$^{-1}$. The tabulated heat of formation of atomic S is 71.9 kJ mol$^{-1}$.

The estimated uncertainty in the thermochemistry is ±0.8 kJ mol$^{-1}$ arising mainly from uncertainty in the CBS extrapolations. This does not include any allowance for correlation effects beyond perturbative quadruples or whether the new $\Delta H_0$ will be accepted generally. We note that in the computational work by Nagy et al.,$^32$ which also involves sulfur species and to which our approach is broadly similar, typical uncertainties in enthalpy were around 2–3 kJ mol$^{-1}$.

A seven-term NASA polynomial fitted to computed thermodynamic functions over 100–3000 K, suitable for use in standard modeling codes, has the following coefficients $a_1$–$a_7$: 2.99418214 × 10$^4$, 9.00692800 × 10$^{-3}$, −7.39316000 × 10$^{-6}$, 2.67928000 × 10$^{-9}$, −3.51975000 × 10$^{-13}$, 2.31452584 × 10$^4$, and 8.59332874 × 10$^9$. With these coefficients, the heat capacity is given by

$$\frac{C_p}{R} = a_1 + a_2 T + a_3 T^2 + a_4 T^3 + a_5 T^4,$$

the entropy is given by

$$\frac{S}{R} = a_1 \ln T + a_2 T + \frac{a_3 T^2}{2} + \frac{a_4 T^3}{3} + \frac{a_5 T^4}{4} + a_7,$$

and the enthalpy, defined as $H = \Delta H_{298.15} + (H_T - H_{298.15})$, is given by

$$H_T = a_1 + \frac{a_2 T}{2} + \frac{a_3 T^2}{3} + \frac{a_4 T^3}{4} + \frac{a_5 T^4}{5} + a_6 T.$$

### B. Kinetics

Figure 1 shows the singlet and triplet potential energy curves for linear N$_2$S as a function of r(N–S). It may be seen that the triplet system is purely repulsive, as expected by analogy with the well-studied N$_2$O system.$^{10}$ The singlet curve shown dissociates to N$_2$ + S. The wave function for these products would require a multi-reference description, but the T1 diagnostic for the singlet state near the crossing is below 0.02 and so CCSD(T) based on a single-reference wave function should be adequate in this region.$^{34}$ The intersection

<table>
<thead>
<tr>
<th>Species</th>
<th>T-Q$^a$</th>
<th>Q-5$^b$</th>
<th>5-6$^c$</th>
<th>T-6$^d$</th>
<th>T-5$^e$</th>
<th>Q-6$^f$</th>
</tr>
</thead>
<tbody>
<tr>
<td>S</td>
<td>−397.67694</td>
<td>−397.67575</td>
<td>−397.67556</td>
<td>−397.67500</td>
<td>−397.67363</td>
<td>−397.67461</td>
</tr>
<tr>
<td>N$_2$S</td>
<td>−507.13445</td>
<td>−507.13070</td>
<td>−507.12852</td>
<td>−507.12758</td>
<td>−507.12305</td>
<td>−507.12479</td>
</tr>
<tr>
<td>$\Delta H_0$</td>
<td>81.53</td>
<td>80.98</td>
<td>80.58</td>
<td>80.23</td>
<td>79.34</td>
<td>79.62</td>
</tr>
</tbody>
</table>

$^a$Two-point extrapolation via Eq. (2).
$^b$Extrapolation via Eq. (4) fitted to four points.
$^c$Three-point extrapolation via Eq. (3).
$^d$Contribution to energy change for reaction (1) in kJ mol$^{-1}$.
of the curves in Fig. 1 approximately indicates the region of intersystem crossing, although the N–N distances are slightly different. Trial and error indicates that the minimum energy crossing point (MECP) occurs near r(N–N) = 1.105 Å and r(N–S) = 2.232 Å, and that both singlet and triplet states at this geometry lie within 0.2 kJ mol\(^{-1}\) of each other and about 72 kJ mol\(^{-1}\) above N\(_2\) + S (\(\ell\)P). The MECP is therefore 144 kJ mol\(^{-1}\) above the v = 0 level of N\(_2\)S. For comparison, in the case of the N\(_2\)O = N\(_2\) + O system the QCISD(T) study of Hwang and Mebel yielded a MECP geometry with r(N–N) = 1.106 Å and r(N–O) = 1.787 Å lying 97 kJ mol\(^{-1}\) above separated N\(_2\) + O (\(\ell\)P).\(^{12}\)

We treat the kinetics of reaction (1) via Rice–Ramsperger–Kassel–Marcus (RRKM) theory. At the low pressure limit the populations of energy levels of N\(_2\)S at the MECP will be controlled by the balance between \(k_1(E)\) and \(k_{-1}(E)\), and in this microcanonical equilibrium the probability of intersystem crossing (which affects both rate processes equally) cancels. The low-pressure limit \(k_{1,0}\) is therefore independent of any assumptions about intersystem crossing,\(^{9,35}\) but does depend critically on the energy of the MECP, and can be treated as a normal reaction via RRKM theory. We will apply Troe’s unimolecular formalism\(^{36}\) to derive \(k_{1,0}\) via

\[
k_{1,0} = Z_{LJ}\beta_c \frac{\rho_{vib}(E_0)RT}{Q_{vib}} \exp \left(-\frac{E_0}{RT}\right) F_{\text{rot}} F_{\text{anh}} F_{\text{rot}} \tag{5}
\]

with energies expressed in J mol\(^{-1}\). The various quantities follow the standard notation. \(E_0\) is the threshold energy for dissociation, i.e., the 0 K bond dissociation enthalpy plus the energy of the MECP relative to the fragments. We first consider application of Troe’s formalism to two isoelectronic analogs of reaction (1), the dissociations of N\(_2\)O and OCS, where experimental data are available.

For OCS our scans of the triplet and singlet surfaces, conducted as described for N\(_2\)S, lead to a MECP at r(C–O) = 1.136 Å and r(C–S) = 2.405 Å, with an energy 35 kJ mol\(^{-1}\) above CO + S(\(\ell\)P). These data imply a threshold energy of \(E_0 = 337\) kJ mol\(^{-1}\) and the ratio of the moments of inertia of the MECP to the molecule of I\(^{1}\)/I\(^{\infty}\) = 1.77. Lennard-Jones (LJ) parameters for the OCS molecule were estimated from its critical properties to be \(\sigma = 4.55\) Å and \(\epsilon/k_B = 289\) K and parameters for Ar were taken from the literature.\(^{37}\) In order to estimate the Lennard-Jones collision rate \(Z_{LJ}\), Evaluation of Eq. (5) with the collision efficiency \(\beta_c\) set to 1 yields the strong collision dissociation rate constant \(k_{\text{disso,0S}}\). This will be higher than that observed because typically several collisions with bath gas molecules are required to transfer sufficient energy to reactant molecules to exceed the dissociation threshold. The collision efficiency can be evaluated as \(k_{\text{exp,disso,0S}}\). Comparison with experimental data for OCS dissociation in Ar bath gas\(^{38}\) shows that \(\beta_c\) decreases from 0.058 at 2000 K to 0.024 at 3000 K. The corresponding average energy transferred per collision \(-\langle\Delta E\rangle\) is derived via the relation\(^{36}\)

\[
\frac{\beta_c}{1 - \beta_c^{1/2}} = \frac{-\langle\Delta E\rangle}{F_{\text{rot}}k_B T}
\]

to obtain \(-\langle\Delta E\rangle\) values around 1 kJ mol\(^{-1}\).

For N\(_2\)O in an N\(_2\) bath gas, Endo et al. derived \(\beta_c = 0.043\) and 0.023 at 900 and 2000 K, respectively, and thus \(-\langle\Delta E\rangle\) = 0.44 and 0.54 kJ mol\(^{-1}\).\(^{39}\) They also suggested that \(N_2\) was a slightly more efficient collider than Ar, by factors of 1.26 and 1.05 at 900 and 2000 K. We therefore expect that the collision efficiency for OCS in N\(_2\) is likely to be similar to that for Ar (in the context of the other uncertainties involved).

For reaction (1) at the low-pressure limit in N\(_2\) bath gas we used \(E_0 = 144\) kJ mol\(^{-1}\) relative to the v = 0 level of N\(_2\)S, and vibrational frequencies of 463, 463, 741, and 2061 cm\(^{-1}\). The Lennard-Jones parameters for N\(_2\)S were assumed to be the same as for the similar OCS molecule, and the parameters for N\(_2\) as a bath gas were taken from Hippler et al.\(^{37}\) We assumed \(-\langle\Delta E\rangle\) provisionally to be independent of temperature and set it equal to 0.5 kJ mol\(^{-1}\). This implies \(\beta_c\) of 0.04 at 1000 K decreasing to 0.02 at 2000 K. The geometry at the MECP yields a moment of inertia 1.60 times larger than for N\(_2\)S, which is used as the ratio I\(^{1}\)/I\(^{\infty}\) to calculate the centrifugal correction \(F_{\text{rot}}\) for this linear system.\(^{36}\) The results from Eq. (5) can be summarized as

\[
k_{1,0} = 7.6 \times 10^{-10} \exp \left(-\frac{126\text{ kJ mol}^{-1}}{RT}\right) \text{cm}^3\text{ molecule}^{-1}\text{s}^{-1}
\]

over 700 – 2000 K.\(^{40}\)

An uncertainty of a factor of 4 arises from uncertainty in the energy transfer parameters and the \(F_{\text{rot}}\) term, but this rough estimate permits an initial evaluation of N\(_2\)S behavior at high temperatures.

Equation (6) is the low-pressure-limiting rate constant which for a small molecule like N\(_2\)S should be applicable over typical pressures. To explore the range of validity some assessment of the high-pressure-limiting kinetics is needed, which are sensitive to the probability of intersystem crossing. We employ the N\(_2\)O analog to set a lower limit to the pressure range where Eq. (6) can reasonably be applied. The experimental high-pressure limit for N\(_2\)O dissociation is \(k_{\text{expt}} = 1.3 \times 10^{13}\) exp(−262 kJ mol\(^{-1}/RT\)) s\(^{-1}\).\(^{3}\) The activation energy \(E_a\) is only 4 kJ mol\(^{-1}\) above the \(E_0\) cited above for the MECP, and the pre-exponential factor is 2–3 orders of magnitude smaller than typical for a simple bond fission reaction.\(^{40}\) This reflects the small probability for intersystem crossing. By analogy, for N\(_2\)S we therefore expect a high-pressure-limiting rate constant \(k_{\text{expt}}\) of \(-\approx 145\) kJ mol\(^{-1}\). Keeping the same pre-exponential factor for the high-pressure limit as for N\(_2\)O decomposition would mean that, at 1000 K and 10 bar of N\(_2\), the predicted first-order dissociation rate of N\(_2\)S is less than 6% of the high-pressure limit.

An alternative estimate of when fall-off effects might become significant for reaction (1) can be made from the first study of N\(_2\)O dissociation at the high-pressure limit by Olschewski et al.\(^{41}\) They showed that the kinetics do not reach the fall-off region until pressures of at least 10 bar. They also directly related \(k_{\text{expt}}\) to the magnitude of the spin-orbit splitting in atomic oxygen. The greater spin-orbit coupling in sulfur as compared to oxygen means that the pre-exponential factor will be greater for N\(_2\)S than for N\(_2\)O at the high-pressure limit, and so Eq. (6) is applicable at even higher pressures than 10 bar. Consistent with this idea, the study by Olschewski
et al.\textsuperscript{42} of CO\textsubscript{2} and CS\textsubscript{2} dissociation provides a pre-exponential factor for $k_\infty$ which is an order of magnitude larger for the sulfur analog vs. the oxygen species.

C. Combustion implications

The analogy between the S + N\textsubscript{2} and O + N\textsubscript{2} reactions suggests that atomic sulfur may conceivably be active in formation of reactive nitrogen intermediates and subsequently nitric oxide under favorable conditions. The importance of S + N\textsubscript{2} as an initiating step in formation of NO depends on the rate of the association reaction, as well as the lifetime of N\textsubscript{2}S and the reactivity of N\textsubscript{2}S towards the radical pool. Both the rate of the S + N\textsubscript{2} association and the lifetime $\tau$ of N\textsubscript{2}S are between the corresponding values for the N\textsubscript{2}O and NNH mechanisms, i.e., $k(O + N_2) < k_{-1} < k(H + N_2)$ and $\tau_{NNH} < \tau_{N_2S} < \tau_{N_2O}$. Thus, in principle, nitrogen fixation via the reverse of reaction (1) is feasible.

To investigate this possibility in more detail, we have carried out chemical kinetic modeling for oxidation of a CO/H\textsubscript{2}/H\textsubscript{2}S mixture at 1800 K and 1 bar in a perfectly stirred reactor under conditions ranging from fuel-lean to fuel-rich. The H\textsubscript{2}S inlet concentration was representative of the level encountered for volatile S in coal combustion, i.e., 1000 ppm, and a CO/H\textsubscript{2} mixture was selected as a fuel to avoid the complexities of hydrocarbon/sulfur interactions. The stirred reactor environment was chosen because this type of reactor augments the radical levels during combustion. The reaction mechanism was drawn from Glarborg,\textsuperscript{43} except for the N\textsubscript{2}S subset, which was estimated in the present work. The following reactions were considered:

\begin{align*}
N_2S + H &\rightarrow N_2 + SH, \quad (7) \\
N_2S + H &\rightarrow NS + NH, \quad (8) \\
N_2S + O &\rightarrow N_2 + SO, \quad (9) \\
N_2S + O &\rightarrow NO + SO, \quad (10) \\
N_2S + OH &\rightarrow N_2 + HOS. \quad (11)
\end{align*}

All of the reactions of N\textsubscript{2}S were assumed to be very fast, with $k_7 = k_9 = k_{10} = 1.5 \times 10^{-10}$ cm\textsuperscript{3} molecule\textsuperscript{-1} s\textsuperscript{-1}, and $k_{11} = 8 \times 10^{-11}$ cm\textsuperscript{3} molecule\textsuperscript{-1} s\textsuperscript{-1}. Reaction (10) would provide a direct pathway from N\textsubscript{2}S to NO, similar to the NNH + O $\rightarrow$ NH + NO process. The rate of reaction (8) was calculated from an estimate of the reverse step, as described below.

Selected modeling predictions are shown in Fig. 2. The calculations indicate that over the range of fuel-air equivalence ratios $0.7 \leq f \leq 1.6$ and temperatures of 1500–1800 K, the formation of NO from the N\textsubscript{2}S mechanism is always well below 0.1 ppm, compared to levels of NO by other mechanisms of 3–12 ppm (1500 K) and 23–39 ppm (1800 K) in the present work. Nitric oxide is formed from N\textsubscript{2}S by reaction with atomic oxygen (reaction (10)). The reason for the low impact of the N\textsubscript{2}S mechanism is that the S-atom concentration is always much lower than those of the O/H radicals.

Under reducing conditions, where [S] peaks, the [H]/[O] ratio is almost 100, and reaction (10) is insignificant compared to reaction (8). Under stoichiometric and lean conditions, where the rates of 7 and 10 become comparable in magnitude, the concentration of atomic S is small, 3–4 orders of magnitude lower than [H] and [O], and negligible amounts of N\textsubscript{2}S are formed. Based on these calculations, the N\textsubscript{2}S mechanism appears to be unimportant for forming NO from N\textsubscript{2}S in most combustion systems. A possible exception could be high temperature industrial processes with high sulfur concentrations, such as combustion of H\textsubscript{2}S or the Claus process.

In addition to the formation through recombination of S with N\textsubscript{2}, N\textsubscript{2}S may be formed from radical-radical reactions. By analogy to steps forming N\textsubscript{2}O, we would expect the reactions between NS and either NH or NCO to be active:

\begin{align*}
NH + NS &\rightarrow N_2S + H, \quad (8') \\
NCO + NS &\rightarrow N_2S + CO. \quad (12)
\end{align*}

As a first estimate these steps would be expected to behave similarly to the analogous reactions involving N\textsubscript{2}O. The NS radical has been detected in sulfur-seeded flames\textsuperscript{44} and reactions (8') and (12) may contribute to the conversion of reaction nitrogen species to N\textsubscript{2}. Observations from a very fuel-rich methane flame\textsuperscript{2b} show that addition of SO\textsubscript{2} accelerates the decay of NO in the post-flame zone. However, despite years of research, the direct interaction of nitrogen and sulfur species in flames is still controversial, and further work is desirable to identify the important steps.

IV. CONCLUSIONS

The enthalpy of formation of singlet N\textsubscript{2}S has been characterized by coupled cluster methods, along with the minimum energy crossing point between singlet and triplet potential energy curves as the N–S distance is increased. The results are used to predict the kinetics of spin-forbidden N\textsubscript{2}S decomposition to N\textsubscript{2} + S(3P) at the low-pressure limit. Modeling of high temperature CO/H\textsubscript{2} oxidation indicates that under stoichiometric and lean conditions the concentration of atomic sulfur is likely to be too small for S + N\textsubscript{2} $\rightarrow$ N\textsubscript{2}S followed...
by O + N₂S → NO + NS to be a significant source of NO, while under fuel-rich conditions the dominant path for N₂S consumption is via reaction with H atoms.

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