brational frequencies from one-photon spectroscopy, and new information can be expected from two-photon spectroscopy. The \( b_2 \) vibrations have been assigned by the following procedure: As the ground electronic state frequencies for \( v_4 \) and \( v_9 \) vibrations are known from the literature, the \( 9^0 \) hot bands (27 830.0 cm\(^{-1}\)) and the \( 9^0 \) hot bands (27 830.0 cm\(^{-1}\)) can be readily assigned in the two-photon spectrum. In Figure 5a,b the relevant parts of the two-photon spectrum are shown. Both the \( 8^0 \) as well as the \( 9^0 \) bands are located on a background resulting from the wings of the rotational envelopes of the strong \( 4^2 \) (27 328.0 cm\(^{-1}\)) and the \( 4^2 \) (27 888.7 cm\(^{-1}\)) transition, respectively. Even though the details of the rotational structure of the \( 8^0 \) and \( 9^0 \) bands cannot be resolved, it is seen that they differ from that of the \( a_1, a_2, \) and \( b_1 \) vibrational bands. Striking features are the sharp cutoff at the blue edge of the band, two small peaks at the red side of the band (marked with arrows), and the lack of a dominating \( R \) branch. No changes in the rotational contour of the non-rotationally symmetric \( A_2 \) two-photon transition are expected when polarization is changed from linear to circular, and indeed, this is detected for these assigned bands. The rotational origins of the hot bands at 27 283.2 cm\(^{-1}\) (\( 8^0 \)) and 27 830.0 cm\(^{-1}\) (\( 9^0 \)) are found from a comparison with the infrared\(^1\) and Raman\(^1,6,17\) data. They are located at the sharp cutoff of the bands, as indicated in Figure 5.

Weak vibronic bands with similar rotational envelopes are found in the cold-band two-photon spectrum at 29 245.4 cm\(^{-1}\) (\( 8^0 \)) and 28 723.6 cm\(^{-1}\) (\( 9^0 \)). From this, we find a value of 871.3 cm\(^{-1}\) for the \( v_4 \) and a value of 349.5 cm\(^{-1}\) for the \( v_9 \) vibrational frequencies in the \( 1^B_1 \) excited electronic state in reasonable agreement with the frequencies calculated by Rolfe et al.\(^20\).

**Summary and Conclusion**

In this work, the low-pressure gas-phase two-photon excitation spectrum of the photodissociating molecule difluorodiazirine, \( \text{F}_2\text{CN}_2 \), was measured under Doppler-limited resolution. On the basis of a detailed analysis of the hot-band region and the analysis of the gross features of the rotational envelopes, nearly all observed transitions have been assigned. In particular, \( b_1 \) vibrational bands can be unambiguously assigned by their striking polarization behavior in linear vs. circularly polarized light.

The bands with \( a_1, b_1, \) and \( b_2 \) vibrational symmetry were discussed in detail. For some of these bands, revised or new assignments are made. The \( v_6 \) vibration, the only vibration of \( a_2 \) symmetry, was not discussed in detail; its previous assignment was corroborated by the two-photon spectroscopy of this work.

Finally, a complete set of vibrational frequencies in the excited electronic \( (1^B_1) \) state of difluorodiazirine is obtained. In particular, the frequencies for \( v_8 \) (1317.7 cm\(^{-1}\)), \( v_9 \) (871.3 cm\(^{-1}\)), and \( v_9 \) (349.5 cm\(^{-1}\)) are found, and new frequency values for \( v_5 \) (785.1 cm\(^{-1}\)) and \( v_5 \) (472.2 cm\(^{-1}\)) are evaluated, which differ considerably from previous values (865.7 cm\(^{-1}\), 332.9 cm\(^{-1}\)) found from a one-photon analysis.\(^3,10,11\)

The ground-state and the excited electronic state frequencies found in this work are listed in Table I together with theoretical values and with previously found values from one-photon and Raman measurements.

It is interesting to note that there is nearly no frequency change \( \delta \) of \( v_4 \) (\( \delta = 15.8 \) cm\(^{-1}\)) and \( v_5 \) (\( \delta = -7.0 \) cm\(^{-1}\)) from the ground to the excited electronic state. \( v_4 \) is a symmetric and \( v_5 \) is an asymmetric CF bond deformation mode. In addition, the frequency of the asymmetric CF stretching mode \( v_6 \) (\( \delta = 69.1 \) cm\(^{-1}\)) increases slightly in the excited electronic state.

By contrast, the vibrations \( v_4 \) and \( v_6 \) leading to a symmetric stretching not only of the CF bond but also of the CN bond (\( v_5 \)) and additionally the NN bond (\( v_9 \)) display a strong decrease (\( \delta = -497.3 \) cm\(^{-1}\)) and a moderate decrease (\( \delta = -162.0 \) cm\(^{-1}\)) in the excited state. A similar behavior is observed for the \( v_4 \) vibration (\( \delta = -219.6 \) cm\(^{-1}\)), which is an asymmetric CN stretching mode.

From these results, it is concluded that there is a tightening of the CF bonds and a loosening of the CN bonds in the electronic excited state. This result is in line with the dissociation pathway of \( \text{F}_2\text{CN}_2 \) leading to \( \text{CF}_2 \) and \( \text{N}_2 \) and may be further confirmed by a detailed analysis of the rotational structure of the various vibronic bands yielding accurate rotational constants. Here, the application of Doppler-free techniques (like Doppler-free two-photon absorption) might be helpful and will be accomplished in forthcoming experiments in this laboratory.

**Acknowledgment.** We are indebted to Prof. E. W. Schlag for his permanent interest and grateful to Dr. E. Riedle for helpful discussions and experimental assistance. Financial support from the Deutsche Forschungsgemeinschaft is gratefully acknowledged. H.S. thanks the Hanns-Seidel-Stiftung for a "Graduierten Stipendium".

**Registry No.** Difluorodiazirine, 693-85-6.

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**Ammonia Activation by \( V^+ \): Electronic and Translational Energy Dependence**

D. E. Clemmer, L. S. Sunderlin, and P. B. Armentrout*†

Department of Chemistry, University of Utah, Salt Lake City, Utah 84112 (Received: May 16, 1989)

The reaction of \( V^+ \) with ammonia is studied as a function of translational energy in a guided ion beam tandem mass spectrometer. The effect of electronic energy is also probed by varying the conditions for forming \( V^+ \). The \( 3^2 \) state of \( V^+ \) is found to be substantially more reactive than the \( 3^1 \) ground and \( 3^1 \) first excited states. The results indicate that reaction occurs primarily through a triplet \( V^+ = \text{NH}_3 \) intermediate. The reactivities of the different electronic states of \( V^+ \) can be explained by using molecular orbital and spin conservation concepts. The thresholds for the two transitions of the endothermic reactions are interpreted to give the 298 K bond energies of \( D^0(V^+ = \text{NH}_3) = 5.18 \pm 0.10, D^0(V^+ = \text{NH}) = 4.30 \pm 0.16, \) and \( D^0(V^+ = \text{N}) = 4.65 \pm 0.06, \) all in electronvolts. The large bond strengths of \( V^+ = \text{NH}_3 \) and \( V^+ = \text{NH} \) indicate that the lone pair electrons on the nitrogen atom are involved in the metal-ligand bond.

**Introduction**

The study of the reactions of atomic transition metal ions with \( \text{H}_2 \) and hydrocarbons has provided insight into the activation of hydrocarbon bonds.\(^1,2\) State-specific studies of the periodic trends in this chemistry have provided a more complete understanding of the electronic requirements for such \( \text{H-H}, \text{C-H}, \) and \( \text{C-C} \) bond activations.\(^1\) Here, we extend this work to examine the activation of the \( \text{N-H} \) bonds of ammonia. The activation of \( \text{N-H} \) bonds

2. State-specific studies of the periodic trends in this chemistry have provided a more complete understanding of the electronic requirements for such \( \text{H-H}, \text{C-H}, \) and \( \text{C-C} \) bond activations.
Ammonia Activation by V⁺

The present study reports guided ion beam studies of the reaction of V⁺ with ammonia. Buckner, Gerd, and Freiser (BGF) have previously examined this system by using Fourier transform ion cyclotron resonance (FTICR) mass spectrometry. They observed the exothermic formation of VNH⁺ as the only reaction at thermal energies. BGF noted that the exothermicity of this reaction means that the V⁻=NH bond strength exceeds D₀(NH2-H₂), which they cited as 4.03 eV. More recent thermochemistry establishes that this limit is actually 4.18 ± 0.01 eV (Table I). BGF also determined an upper limit to this bond energy by observing the exothermic reaction 1. On the basis of D₀(V⁺-O)

\[
\text{VNH}^+ + \text{H}_2 \text{O} \rightarrow \text{VO}^+ + \text{NH}_3
\]

It was found that V⁺ with ammonia over an extended kinetic energy range, such that we study the endothermic as well as the exothermic reactions. This permits a detailed dynamic study of the interaction of V⁺ with ammonia and allows a comprehensive evaluation of the thermochemistry of V⁺-NH₄⁺ (n = 0–2). In addition, we compare these results with the previous examination by Aristov and Armentrout (AA) of the state-specific reactivity of V⁺ with methane. Because methane is isoelectronic with ammonia (in the sense that the central heavy atom has the same number of valence electrons with the same sp³ hybridization), this study provides an excellent reference point for understanding the present results.

Results of the methane experiment show that electronic energy from excited-state vanadium ions couples efficiently to the reaction coordinates of all product channels. The reactivity of the a1D state was found to be over 2 orders of magnitude greater than that of the a2D ground state. Bond strengths of the product molecules VCH⁺⁺, VCH⁺, and V⁺ were reported. Experimental results led to a proposed reaction mechanism in which the first step involves oxidative addition of the C-H bond at the metal center to form H-V⁺-CH₃. At low energies, H₂ is eliminated via a four-center transition state to form VCH⁺. At higher energies, cleavage of the V⁻=O or V⁻=C bond of the intermediate leads to production of VCH⁺⁺ or VH⁺, respectively. V⁺ is formed primarily by loss of H₂ from the VCH⁺⁺ product but also by loss of H from VCH⁺.

The major difference between CH₄ and NH₃ is the lone pair of electrons present on ammonia. We examine the effect that this lone pair has upon the cross-section magnitudes, ionic thermochemistry, and reaction mechanism by comparing this system to the methane system. The information gained may thus provide insight into the interaction of transition metals with such electrons and thereby improve the coordination chemist’s view of metal-ligand bonding.

Experimental Section

General. A complete description of the apparatus and experimental procedures is given elsewhere. Briefly, the apparatus comprises three differentially pumped vacuum chambers. In the first chamber, ions are produced as described below. The resulting ions are extracted, accelerated, and focused into a magnetic sector mass analyzer for mass analysis. In the second vacuum chamber, the mass-selected ions are decelerated to a desired kinetic energy and focused into an octopole ion guide. Radio frequency electric fields in the guide create a radial potential well which traps ions over the mass range studied. The velocity of the ions parallel to the axis of the guide is unchanged. The octopole passes through a static gas cell into which ammonia can be introduced. Pressures are maintained at a sufficiently low level (less than 0.2 mTorr) that multiple ion–molecule collisions are improbable. Product and unreacted beam ions are contained in the guide until they drift out of the gas cell. The ions are then extracted and focused into the third vacuum chamber which contains a quadrupole mass filter for product mass analysis. Ions are detected with a secondary electron ionization detector and pulse-counting electronics. Raw ion intensities are converted to absolute cross sections as described previously. The accuracy of our absolute cross sections is estimated to be ±20%. Uncertainties at low-cross-section values are generally about ±10⁻¹⁰ cm², primarily because of random counting noise (typically ±10 counts/s). Translational energies in the laboratory frame of reference are related to energies in the center of mass (CM) frame by E_CM = E_lab/(M+m), where M and m are the masses of the incident ion and neutral reactant, respectively. For these experiments, V⁺ (99.76% natural abundance) was used. The absolute energy scale and the ion kinetic energy distribution are determined by using the octopole beam guide as a retardation potential analyzer. Uncertainties in the absolute energy scale are ±0.05 eV lab, and the fwhm of the energy distribution is 0.6 eV lab. Below ~0.6 eV

<table>
<thead>
<tr>
<th>TABLE I: Heats of Formation and Bond Strengths for NH₄ Species (298 K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>neutral ΔH°, eV</td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td>H</td>
</tr>
<tr>
<td>N</td>
</tr>
<tr>
<td>NH</td>
</tr>
<tr>
<td>NH₂</td>
</tr>
<tr>
<td>NH₃</td>
</tr>
</tbody>
</table>

*a All values except where noted are from ref 15. Uncertainties are in parentheses. bDerived from ΔH° results for NH and NH₂ in: Anderson, W. R. J. Phys. Chem. 1989, 93, 530-536.

<p>|TABLE II: V⁺ Beam State Populations |
|---|---|---|</p>
<table>
<thead>
<tr>
<th>electron config</th>
<th>eV</th>
<th>% population</th>
</tr>
</thead>
<tbody>
<tr>
<td>a1D 3d⁴</td>
<td>0.026</td>
<td>84.0 (1.4)</td>
</tr>
<tr>
<td>a1F 4s³d⁳</td>
<td>0.36</td>
<td>15.9 (1.4)</td>
</tr>
<tr>
<td>a1F 4s³d³</td>
<td>1.10</td>
<td>0.116 (33)</td>
</tr>
<tr>
<td>a1P 3d⁴</td>
<td>1.45</td>
<td>0.006</td>
</tr>
<tr>
<td>a1P 3d⁴</td>
<td>1.57</td>
<td>0.012</td>
</tr>
<tr>
<td>b1F 4s³d³</td>
<td>1.69</td>
<td>0.004</td>
</tr>
<tr>
<td>b1P 3d⁴</td>
<td>1.69</td>
<td>0.002</td>
</tr>
<tr>
<td>b1G 3d⁴</td>
<td>1.81</td>
<td>0.002</td>
</tr>
<tr>
<td>Σtriples</td>
<td>&lt;0.001</td>
<td></td>
</tr>
</tbody>
</table>

*Energies are averaged over J levels taken from ref 36. Maxwell-Boltzmann distribution. Uncertainties due to the ±100 K spread in filament temperatures are in parentheses. Estimated value from VCH⁺⁺ channel from ref 16.

(14) We use ΔH°; (H₂O) = 59.55 kcal/mol and ΔH°; (H₂) = 57.80 kcal/mol.
lab (0.15 eV CM), the energies are corrected for truncation of the ion beam energy distribution as described previously. The data obtained in this experiment are broadened by the ion energy spread and thermal motion of the neutral gas. The second effect, referred to as Doppler broadening, has a width (in eV) in the CM frame of 0.47E1/2 for the reaction of V+ with NH3. When model cross sections are compared to experimental data, the calculated cross sections are convoluted with both sources of experimental energy broadening as described previously.

Ion Sources. Vanadium ions are produced in two sources in these experiments. The primary source is a surface ionization (SI) source. Here, gaseous VOC13 (Alfa 99.99%) is directed at a resistively heated rhenium filament where decomposition of VOC13 and ionization of the resultant vanadium atoms takes place. The temperature of the filament is calibrated by optical pyrometry measurements and has an absolute uncertainty of ±100 K. It is generally assumed that a Maxwellian distribution of translational energy in the reactants is used. In this study, we have utilized eq 6.20,22 we have utilized eq 6.20,22,26 20 Table II gives these populations for V+ at the filament temperatures used in these experiments.

The second source of ions used in these studies is an electron impact (EI) source. The electrons ionize and dissociate VOC13 vapor, leaving V+ and other ions. The state populations are not characterized by a Maxwellian distribution since ionization is dominated by vibrational processes. Since the appearance potential of V+ from VOC13 is 26.8 ± 0.4 eV,21 it is possible to form significant percentages of excited-state ions at the electron energies of 30, 50, and 70 eV used in these experiments. This has been verified in previous experiments.20,22 The populations measured are included in Table II.

An electron impact/drift cell (DC) source26 was also used in an attempt to electronically cool V+ by collisions with methane. Although conditions in this source were such that the ions underwent over 1000 collisions with methane, noticeable contributions to the cross section from ions in excited electronic states were still observed. The neutral gas used in this experiment was anhydrous ammonia (Matheson 99.99%). Before usage, three or more freeze–pump–thaw cycles with liquid nitrogen were done to remove volatile impurities.

Thermochemical Analyses. Theory23 and experiment6,20,22,26 indicate that cross sections can be parametrized in the threshold region by eq 2, where \( \sigma_0(E) \) is a scaling factor, \( E \) is the relative translational energy of the reactants, \( E_T \) is the threshold energy, and \( m \) and \( n \) are adjustable parameters. As in previous studies,16,20,22,26 we have utilized eq 2 with \( m = 1 \). This form is expected to be the most appropriate for translational energy in the threshold region in the present system as well. Furthermore, we make no correction for the energy available in internal modes of the neutral reactants and the products formed at the threshold of an endothermic reaction are characterized by a temperature of 298 K in all degrees of freedom.

Figure 1. Variation of product cross sections for reaction of ammonia with V+ produced by SI as a function of translational energy in the center-of-mass frame (lower scale) and laboratory frame (upper scale). The solid line is the sum of the cross sections for all products. The dashed line is the sum of \( \sigma(VN^+) \) and \( \sigma(VNH_2^+) \).

linear least-squares analysis to give the best fit to the data. The cited threshold is the average of the analyses for all data sets. Conservative error limits for \( E_T \) are calculated from the range in these threshold values for different data sets and the error in the absolute energy scale (0.02 eV CM). These thresholds depend strongly upon the electronic state of the reactant ion and are converted to the threshold for the ground state, \( E_0 = E_T + E_{el} \) where \( E_{el} \) is the average electronic energy of the reactants. Analysis can also be accomplished by explicitly including the various states in eq 2. Such analyses yield \( E_0 \) values which are consistent within 0.03 eV with the values derived by correcting \( E_T \) with \( E_{el} \).

For some data channels it is necessary to use a modified form of eq 2 which accounts for cross sections which decline due to dissociation of the product ion at higher energies. We have previously outlined such a model which makes a simple statistical assumption within the walls of previous studies.23 This empirical model defines \( P_{el} \), the probability of dissociation of the product ion, as a function of \( E_{el} \), the energy at which dissociation begins, and \( p \), which is related to the number of internal modes in the transition state. Thus for \( E < E_{el} P_{el} \) is zero, and for \( E > E_{el} P_{el} \) asymptotically approaches 1. In this study, \( E_0 \) is optimized to best fit the data. The parameter \( p \) (which is limited to integer values) is treated as a variable but the optimum value in this study is always 3.

Reaction threshold energies are converted to thermochemical values of interest by assuming that \( E_T \) represents the enthalpy difference between reactants and products. This assumes that there are no activation barriers in excess of the endothermicity. This assumption is generally true for ion-molecule reactions in this system as well, and we have explicitly tested a number of times,26 and appears to be a reasonable one in the present system as well. Furthermore, we implicitly assume that the neutral reactants and the products formed at the threshold of an endothermic reaction are characterized by a temperature of 298 K in all degrees of freedom. Thus, we make no correction for the energy available in internal modes of the neutral reactant.

Results

Four products, formed in reactions 3–6, are seen in the reaction of V+ with NH3. Figure 1 shows the cross sections for these...
products when V$^+$ is produced at a surface ionization (SI) filament temperature of 1950 K. (Comparable results are obtained for reaction with ND$_3$.) Below ~1.6 eV, reaction 5 dominates the reactivity. The fact that this cross section increases with decreasing energy to as low an energy as we can measure indicates that this reaction is exothermic, in agreement with BGF.$^3$ To be exothermic, formation of VNH$^+$ must be accompanied by production of the stable H$_2$ molecule. This is confirmed by noting that the VNH$^+$ cross section also exhibits an endothermic feature with a threshold of ~0.8 eV, but there is also a small exothermic component to this cross section. The endothermic feature peaks at about 2.8 eV. This is well below the energy needed to dissociate to V$^+$ + H$_2$ (4.69 eV, Table I), which is marked in Figure 1. The early peak could be due to dissociation to VN$^+$ + H$_2$ which has a threshold at about this energy; however, as shown by the sum of $\sigma$(VNH$_2^+$) and $\sigma$(VN$^+$) (Figure 1) the magnitude of the VNH$^+$ cross section is too small to account completely for the decline in VNH$_2^+$. The only remaining explanation is that the VH$^+$ channel, reaction 3, is influencing the VNH$_2^+$ cross section. This is consistent with the smooth appearance of the total cross section and is an indication that reactions 3 and 6 compete directly. At the highest energies, reaction 3 dominates the reactivity. Reaction 4 is always a minor channel. One reason is that the VN$^+$ product must be formed via decomposition of other products, reactions 8 and 9. Comparison of the magnitudes of $\sigma$(VNH$_2^+$)

\[
\text{VNH}_2^+ \rightarrow \text{VN}^+ + \text{H}_2
\]  

(8)

and $\sigma$(VNH$^+$) to $\sigma$(VN$^+$), Figure 1, suggest that $\sigma$(VNH$_2^+$) is too small to account for the magnitude of $\sigma$(VN$^+$). Further, we find that the sum of the VN$^+$ and VNH$_2^+$ cross sections (shown in Figure 1) falls off smoothly, consistent with VNH$_2^+$ being the major contributor to VN$^+$ formation via reaction 8. The change in slope in the summed cross section curve at 4.7 eV is due to dissociation of VNH$_2^+$ to V$^+$ + NH$_3$, which can begin at $D^e$(NH$_2$-H) = 4.69 eV. This dissociation channel is also reflected in the VN$^+$ cross section, an observation which is further evidence for the parentage of VN$^+$. Similar behavior can also be seen in the higher energy feature of VNH$^+$. This is evidence that reaction 7 proceeds via decomposition of VNH$_2^+$ to form VNH$^+$ + H (the only plausible pathway).

As mentioned above, the electronic state of V$^+$ can greatly affect the reaction cross section. In order to discuss these effects quantitatively, we examine SI and electron impact (EI) data for each individual product channel and derive state-specific cross sections when possible. Reaction thresholds are determined for each product channel and then used in the Discussion section to derive bond strengths for the product molecules.

\[\text{VH}^+ \]  

The most efficient process above ~2.5 eV in the V$^+$ + NH$_3$ reaction is the production of VH$^+$. The SI cross sections for this product are independent of the filament temperature at which V$^+$ is produced. Analysis of these SI data sets, Table III, give an average threshold of $E_T = 2.52 \pm 0.09$ eV. When corrected for $E_d = 0.09 \pm 0.02$ eV, we find $E_d = 2.61 \pm 0.09$ eV. Given the value of $D^e$(NH$_2$-H) listed in Table I, this yields a bond energy for V$^+$-H of 2.08 \pm 0.09 eV, which is in excellent agreement with our previously reported VH$^+$ bond energy of 2.09 eV.

### Table III: Summary of Parameters of Eq 2 Used To Fit Threshold

<table>
<thead>
<tr>
<th>product</th>
<th>source$^a$</th>
<th>$E_T$, eV</th>
<th>$\sigma_0$, eV</th>
<th>$\sigma$, eV</th>
</tr>
</thead>
<tbody>
<tr>
<td>VH$^+$</td>
<td>SI 1950 K</td>
<td>2.57 (0.04)</td>
<td>0.36 (0.02)</td>
<td>1.0 (0.1)</td>
</tr>
<tr>
<td>SI 2200 K</td>
<td>2.46 (0.03)</td>
<td>0.44 (0.02)</td>
<td>1.0 (0.1)</td>
<td></td>
</tr>
<tr>
<td>EI 30 eV</td>
<td>1.63 (0.08)</td>
<td>3.35 (0.25)</td>
<td>1.1 (0.1)</td>
<td></td>
</tr>
<tr>
<td>VN$^+$</td>
<td>X(SI)</td>
<td>2.91 (0.06)</td>
<td>0.066 (0.006)</td>
<td>1.0 (0.1)</td>
</tr>
<tr>
<td>EI 30 eV</td>
<td>1.52 (0.12)</td>
<td>0.46 (0.05)</td>
<td>1.4 (0.4)</td>
<td></td>
</tr>
<tr>
<td>VNH$^+$</td>
<td>X(SI)</td>
<td>4.33 (0.16)</td>
<td>0.026 (0.003)</td>
<td>1.0</td>
</tr>
<tr>
<td>VNH$_2^+$</td>
<td>X(SI)</td>
<td>1.47 (0.06)</td>
<td>0.074 (0.005)</td>
<td>1.6 (0.2)</td>
</tr>
</tbody>
</table>

$^a$This refers to the V$^+$ source used to obtain the raw data analyzed. In cases where the $X$ or $a^3F$ state is referred to, the text provides a complete description of the derivation of the cross section analyzed.

Figure 2. Kinetic energy dependence of $\sigma$(VH$^+$) formed by reaction of ammonia with V$^+$ produced by SI at 1950 K (●), by EI at 30 eV (●), by EI at 50 eV (○), and by EI at 70 eV (□) as a function of translational energy in the laboratory frame (upper axis) and the center-of-mass frame (lower axis).

The agreement implies that this threshold is not subject to obvious systematic errors, such as misidentification of the reactive state(s) or activation barriers in excess of the endothermicity.

Figure 2 compares the VH$^+$ cross section obtained from V$^+$ formed in the SI source to those obtained for V$^+$ formed with the EI source at electron energies of 30, 50, and 70 eV. The endothermicities, shapes, and magnitudes of the cross sections are clearly very dependent upon the source of V$^+$. The 30-eV cross section is about an order of magnitude greater than the SI cross sections and is nonzero at the lowest energies we can measure. Analysis of the main endothermic feature of this cross section using eq 2 and the fitting parameters given in Table III yields a threshold of $E_T = 1.63 \pm 0.08$ eV. Thus, $E_d = E_d - E_T$(30 eV) = 0.98 \pm 0.12 eV, a value which roughly correlates to the excitation energy for the $a^3F$ state of 1.10 eV (Table II). Thus, we conclude that $\sigma$(VH$^+$) from the 30-eV beam is dominated by V$^+$($a^3F$). The notable contribution to the 30-eV cross section at energies below 1.63 eV indicates that states higher in energy than the $a^3F$ are also populated. AA have previously estimated the $a^3F$ population of the 30-eV beam to be 9 \pm 3%. This estimate suggests that the cross section for VH$^+$($a^3F$) is about a factor of 10 greater than the endothermic feature of the 30-eV cross section shown in Figure 2.

Increasing the electron energy from 30 to 50 eV significantly increases the low-energy cross section for VH$^+$ formation. This is clearly an exothermic process which indicates that ions with electronic energies at least 2.5 eV above the ground state are contributing to the 30-eV VH$^+$ cross section. At higher kinetic energies, the cross section has an apparent onset and magnitude.
which matches the 30-eV data. Thus, this endothermic portion of the cross section is dominated by $V^+(a_{3F})$.

Upon increasing the electron energy from 50 to 70 eV, we observe that the magnitude of the exothermic portion of the cross section decreases. The endothermic portion has the same magnitude and apparent onset as the 30-eV data and thus corresponds to $a_{5D}$-state contribution. The decrease in the exothermic pathway indicates that the states contributing to that process have been depopulated.

At high kinetic energies, $\sigma(VH^+)$ could decline due to the decomposition to $V^+ + H$ which can begin at 4.69 eV (Table I). Since $\sigma(VH^+)$ does not decline at higher kinetic energies, the excess energy must be contained in either kinetic energy or internal modes of the NH$_2$ product. This is typical behavior for metal hydride ion products formed in reactions with alkanes.16,20,23

$VN^+$. The most endothermic process in the reaction of $V^+$ with ammonia is the formation of vanadium nitride ions, reaction 4. Figure 3 shows the 1950 and 2200 K SI cross sections for this reaction. At an energy near the threshold, 2.4 eV, the cross section is enhanced by a factor of ~2.3 when the SI filament temperature is increased from 1950 to 2200 K. The only plausible explanation for this enhancement is the increase in the population of the low-lying triplet states. Indeed, Table II shows that the populations of the $a_{5F}$ and higher lying states increase by a factor of 2.1 as the filament temperature increases from 1950 to 2200 K. At higher kinetic energies, the magnitude of the cross section changes little with filament temperature, indicating that this reactivity is dominated by the $a_{3D}$- and $a_{5F}$-state ions.

Figure 3 also shows $\sigma(VN^+)$ formed by $V^+$ beams produced by EI with electron energies of 30 and 50 eV. The EI data have a greater magnitude than the SI data and show thresholds which have been shifted to lower energies. Specifically, the 30-eV cross section yields a threshold of 1.52 ± 0.12 eV when fit with the parameters given in Table III, and peaks at about 3.4 eV. This ~1-eV shift in cross section peak position between the SI and 30-eV EI data correlates roughly to $E_D(a_{5F})$. Thus, the $a_{5F}$ population of the 30-eV beam is apparently responsible for the peak position of the $VN^+$ cross section.

Upon increasing the EI electron energy from 30 to 50 eV, the magnitude of $\sigma(VN^+)$ is increased, and an exothermic as well as endothermic pathway to $VN^+$ formation is observed. The increase in magnitude again indicates that excited states make up an even greater portion of the 50-eV beam than the 30-eV beam. The exothermic pathway indicates that highly excited $V^+$ ions contribute to $VN^+$ formation. Based on the 1.52-eV threshold for the 30-eV EI data, these ions must have at least 2.6 eV (i.e. $E_D(a_{3F})$ internal excitation. The 70-eV EI data also show an exothermic as well as an endothermic process. The magnitudes of both features are similar to the 50-eV data.

Analysis of the $VN^+$ Cross Sections. Cross sections for production of $VN^+$ by the $a_{3D}$ and $a_{5F}$ states are derived by taking the 1950 and 2200 K SI data along with the populations of the excited triplet states given in Table II and extrapolating to a contribution of 0.0% of these states. This leaves a cross section (shown in Figure 3), due to reaction of only $a_{3F}$ and $a_{5F}$ ions, which we will refer to as the $X$ cross section since the relative reactivities of these two states cannot be unequivocally determined. This $X$ cross section can then be analyzed with eq 2 and the parameters in Table III and $E_D = 4.6$ eV. After correcting for $E_D(X) = 0.07 ± 0.02$ eV, we obtain $E_D(a_{3F}) = 2.98 ± 0.06$ eV. By using the same values of $n$ and $m$ as for $\sigma(X)$ and adjusting the threshold by $E_D(a_{5F})$, we can derive a model for the $a_{5F}$ state. When added to the quintet-state model after appropriate scaling, Table II, both SI data sets are accurately reproduced.

As noted above, analysis of the 30-eV EI data yielded $E_D = 1.52$ eV. This suggests that the average value of $E_D$ is 1.46 ± 0.08 eV, demonstrating that states above the $a_{5F}$ are contributing to this cross section. The $a_{5F}$ state is probably significant since $E_D(a_{5F}) = 1.45$ eV, Table II. By combining appropriate amounts of model cross sections for the $a_{5F}$ and $a_{5D}$ states (again using the same values of $n$ and $m$ as for $\sigma(X)$ and adjusting the threshold by $E_D(a_{3F})$), the 30-eV data can also be reproduced.
Ammonia Activation by V⁺

Figure 5. Kinetic energy dependence of $\sigma(VNH\phantom{+}^+)$ formed by reaction of ammonia with V⁺ produced by SI at 1950 K (○), SI at 2200 K (△), and EL at 30 eV (●), and EL at 50 eV (□) as a function of translational energy in the laboratory frame (upper axis) and the center-of-mass frame (lower axis). The dashed line and double-dash line show the contributions of the a3F and higher lying states, respectively, to the 30-eV cross section. The arrow at 3.59 eV shows the bond energy $D^b_0(\text{H}_2\text{N}\text{-H}) - E_2(a^3F)$. Contributions to the SI data from the $^3X(\uparrow)$ and $^6\text{O}(\uparrow)$ (dash-dot line) are also shown.

thermic feature of $\sigma(VNH\phantom{+}^+)$ observed in the SI data is less prominent in the EL data. This is due to the large magnitude increase in the exothermic EI cross section which masks the smaller decomposition pathway, reaction 7.

Analysis of the VNH⁺ Cross Sections. In order to derive a cross section representative of the a3D and a3F states, we again extrapolate the SI data to a contribution of 0.0% of the excited triplet states. This X cross section is shown in Figure 4. For this large energy channel, the data is sufficiently good that we can also extrapolate to 100%, thus generating a cross section for the triplet states, X, also shown in Figure 4.29 The validity of this extrapolation is supported by comparison of the 30-eV data to this latter extrapolated cross section. This comparison shows that the shapes are nearly identical, consistent with expectations based on the reaction channels discussed above. Also, the 30-eV data has a magnitude which is 24 ± 5% of the extrapolated X cross section. Thus, the population of excited triplet states in the 30 eV beam is 24 ± 5%, consistent with the estimate given by AA, 21 ± 5% (Table II).

The X cross section shows that reaction 5 is exothermic with a small endothermic feature at high energies. This establishes that the reaction of the V⁺(a3F) state is exothermic such that $D^b_0(V^+\text{-NH}) \leq 3.81$ eV (4.17–0.36). A more precise value of this bond energy can be obtained by analyzing the cross section for reaction 7, the endothermic feature observed in the SI data for VNH⁺. We arrive at the threshold for this feature by the following method. The exothermic portion of the X cross section (and also the raw SI data) was fit in the energy region below the endothermic feature (between ~2.8 and 3.9 eV) with a power law and this was subtracted from the cross section, leaving only the endothermic feature. This feature was then analyzed with eq 2 by using the parameters given in Table III. The threshold for this feature, after correction for $E_D$, yields $E_D = 4.40 \pm 0.16$ eV. Since the bond energy of H₂ is 4.52 eV, this value is consistent with the exothermicity of reaction 5 for V⁺(a3F) and establishes that reaction of a3D ground-state ions is also exothermic.

$V\text{NH}^+$. Figure 5 shows $V\text{NH}^+$ cross sections from V⁺ produced by the SI source with filament temperatures of 1950 and 2200 K. Clearly, the exothermic portions of these cross sections depend on temperature while the endothermic feature, which peaks at ~1.8 eV, does not. At 1 eV, the 1950 and 2200 K cross sections have magnitudes of about 0.005 and 0.009 Å², respectively, a ratio of 1.0:1.8. Table II shows that the fraction of the a2D state decreases as the temperature is increased, while the a3F increases by a factor of only 1.2. Thus, the differences between the 1950 and 2200 K data must be due largely to the ~10⁻³ fraction of the beam in excited triplet states. Table II shows that the a3F population increases by a factor of 2.0, while higher lying states increase by a factor of 2.7. Thus, the dominant contribution to the cross section in this energy region must come from the a3F. The higher energy states may also contribute to the exothermic cross section but cannot quantitatively account for the temperature dependence.

Also shown in Figure 5 are the $V\text{NH}^+$ cross sections obtained by using the EI source with electron energies of 30 and 50 eV. As in the other channels, these cross sections show an increase in magnitude when compared with the SI data. The 30-eV cross section peaks at ~1.8 eV, about 1 eV lower than the peak in the SI data. This 1-eV shift in peak position correlates well to $E_D(a^3F)$ and suggests that the endothermic portion of the 30-eV data is due to reaction of the a3F state. Therefore, the exothermic feature is presumed to be due to reaction of a3F or higher states.

Increasing the electron energy from 30 to 50 eV increases the low-energy exothermic feature of this cross section by about an order of magnitude. This helps confirm the identification of this feature as coming from a3F states or higher since AA made a similar state identification by observing similar effects in going from 30 to 50 eV EI conditions for the reaction $V^+ + \text{H}_2 \rightarrow \text{VCH}_2^+ + \text{H}_2$ (a reaction which has similar energetics to reaction 6). A curious feature of the 50-eV data is that this change in electron energy also gives rise to enhanced $V\text{NH}_2^+$ production at high kinetic energies (>3 eV). The increase at high energies does not correspond to either the SI or 30-eV EI cross-section shape and must therefore be due to a higher energy state(s).

Increasing the electron energy from 50 to 70 eV has little effect on the magnitude of the low-energy feature of the cross section but does increase the very high energy feature to a cross section of 0.4 Å² at 6 eV. This increase is consistent with highly excited states accounting for this feature. Since these states appear to react mostly efficiently at kinetic energies above ~3 eV, this suggests that these states either exhibit an activation barrier, or produce excited $V\text{NH}_2^+$ and $\text{H}_2$ products in an endothermic reaction.

Analysis of the VNH⁺ Cross Sections. In order to obtain $E_D$ and elucidate a V⁺–NH₂ bond strength, we need to understand how the different electronic states of V⁺ contribute to reaction 6. Since the 30- and 50-eV data sets have different amounts of these states accounting for this feature, since these states appear to react mostly efficiently at kinetic energies above ~3 eV, this suggests that these states either exhibit an activation barrier, or produce excited $V\text{NH}_2^+$ and $\text{H}_2$ products in an endothermic reaction.

ED = 0.38 ± 0.06 eV and $E_D = 1.9$ eV. Assuming that the a3F state is the only contributor to this derived endothermic feature, the cross section $E_D > 1.48$ eV react exothermically. This includes all higher lying states with the possible exception of V⁺(a3P) (Table II). If the a3P reaction is endothermic, our analysis would lead to a higher value for $E_D$ depending on the relative contributions of the a3F and a3P states to the endothermic feature. We have verified that these changes in $E_D$ are small as long as the a3F state is the dominant contributor to the endothermic feature (which seems likely based on results for other product channels). Analysis of the SI cross sections (below) also helps confirm this assumption.

(29) Technically, there may be contributions from the a3P state in this cross section. It is likely that such contributions are small since the reactivity of the quintet states is much smaller than the triplet states.
State-Specific Cross Sections. For each of the reactions discussed above, it was possible to derive a cross section for the a^3D and a^3F states and for excited states, usually the a^3F state. The former cross section, which we will term X, is shown in Figure 6. These cross sections are basically those of the SI data with small contributions from excited states removed. Thus, the absolute magnitudes and shapes of these cross sections are in little doubt. For the excited states, as also shown in Figure 6, the absolute cross sections are less certain but are certainly accurate to within a factor of ~3. These cross sections were derived as follows. The VH+ cross section was derived by taking the model cross section for the 30-eV EI data and correcting the absolute magnitude by using AA's estimate for the population of the a^3F state (Table II). For the VNH+ and VNH2+ cross sections, the models used to reproduce the a^3F contributions to the SI cross sections were scaled by the populations in Table II to obtain absolute a^3F cross sections. A check on the internal consistency of this procedure is to use this absolute a^3F cross section to estimate the population of this state in the 30-eV EI data. We find that this comparison suggests a ~13% population for (VNH2+) and ~10% for (VNH+), in good agreement with AA's value of 9 ± 3%. As a cautionary note, we find that if the 50-eV EI data is similarly used, the absolute cross section for (VNH2+) derived is about 3 times larger than the one shown. This is probably an accurate measure of the absolute uncertainties in these derived cross sections. In the case of V+(a^3P), the cross section shown includes a model which accounts for the observed decline in the cross section at high energies due to dissociation of VNH2+ to V* + NH2, beginning at 3.59 eV = D*(NH2-H) - E_d(a^3P). Finally, for the VNH+ cross section, we cannot extract a cross section due solely to reaction of V+(a^3F) since all states react exothermically to form VNH+. Therefore, we have used a sum of the triplet states, as discussed above, and extrapolated to probe (VNH2+). As shown in Figure 4, this manipulation leads to fairly large scatter, although the shape of the cross section is equivalent to that of the 30-eV EI data. Therefore, Figure 6 shows this latter data scaled to the appropriate absolute magnitude.

The most striking feature of Figure 6 is the large increase in the magnitude of (V+) when compared with (VNH+). On the basis of the VN+ and VNH2+ cross sections, we estimate that V+(a^3F) is more reactive than V+(a^3P) by factors of 110 and 170, respectively. The VNH+ channel shows that the reactivity of V+(a^3P) is greater than that of V+(a^3F) by a factor of ~160. This enhancement in reactivity is comparable to the effect observed by AA for the dehydrogenation reaction of methane by these states of V+

Figure 6 also shows that the competition between the reaction channels discussed above for the 1950 K SI data, which is primarily due to reaction of V+(a^3P), is also apparent for V+(a^3X). Specifically, it can be seen that the peak in (VNH2+) occurs near the threshold for (VNH2+) for both states. The sizes of (VNH+) and the high-energy component of (VNH2+) relative to (VNH2+) are similar for both states. In all cases, the thresholds and peaks of the cross sections are energy shifted by slightly more than 1 eV = E_d(a^3P). This similarity in behavior indicates that the reaction mechanisms of both quintet and triplet states are the same.

Reaction Efficiency. Comparison of the a^3X cross section with a comparison indicates that these states react with ~1% efficiency at our lowest energies (~0.1 eV). By averaging our experimental cross sections over a Maxwell–Boltzmann distribution, we can determine a thermal rate constant which is 9.8 × 10^12 cm^3/s, 1% of that of the total collision rate. This is somewhat lower than the 5% efficiency value given by BGF and suggests that they have excited-state V* contributing to their observed reaction rate. This is a reasonable possibility since BGF produced their vanadium ions by laser desorption/ionization.

The a^3X cross section is comparable to a comparison at the lowest energies (this is most easily seen in Figure 4), which suggests that these
Ammonia Activation by V* 

Excited states react with near unit efficiency. However, ammonia has an appreciable dipole moment (1.47 D) such that the collision frequency should actually be higher than predicted by σ₁. A reasonable upper limit to the true collision cross section is easily calculated for the locked dipole model, \( \sigma_{LD} \). We find that, below 1.0 eV, the \( J^2 \) cross section behaves as \( \sim 0.25(\sigma_{LD}) \).

Discussion

Thermochemistry. Bond energies for \( V^+\text{-H} \) (2.08 ± 0.09 eV), \( V^+\text{-N} \) (4.65 ± 0.06 eV), \( V^+\text{-NH} \) (4.40 ± 0.16 eV), and \( V^+\text{-NH}_2 \) (3.18 ± 0.10 eV) are derived from the ammonia thermochemistry listed in Table II and \( E_0 \) values for reactions 3, 4, 7, and 6 (2.61 ± 0.09, 2.98 ± 0.06, 4.40 ± 0.16, and 1.51 ± 0.10 eV, respectively). The \( V^+ \) bond energy determined here agrees well with a previous value, 2.09 ± 0.06 eV.\(^{22} \) Our \( VN^+ \) bond energy is considerably higher than the 3.78 eV value calculated by Kunze and Harrison; however, these authors note that their absolute bond energies are generally low by about 25%. Correcting our result by this factor provides an estimated bond energy of \( \sim 4.7 \) eV in good agreement with our experimental result. Our \( VN^+ \) value also agrees with the limits of 4.17 eV to 5.17 eV derived from BGF’s work\(^{3} \) as described in the Introduction, as well as their stated result of 4.38 ± 0.30 eV.

We can also compare our value for \( D^0(VN^+) \) = 4.65 ± 0.06 eV to literature values of \( D^0(VN) \) = 4.95 ± 0.09 eV and 4.82 ± 0.22 eV.\(^{35} \) Using these bond strengths and the ionization potential of vanadium, IP(V) = 6.740 ± 0.002 eV,\(^{26} \) eq 10 can be used to calculate IP(VN) = 7.04 ± 0.11 or 6.91 ± 0.23 eV.

\[
IP(VN) = D^0(VN) + IP(V) - D^0(VN^+) \quad (10)
\]

in agreement with the rough experimental value of 8 ± 1 eV.\(^{34} \) Since the IPs of V and VN are similar, ionization of VN probably involves the removal of an electron from a nonbonding metal orbital.

The bond energies \( D^0(VN^+\text{-H}) = 3.10 ± 0.17 \) eV and \( D^0(VN^+\text{-H}^+) = 2.88 ± 0.19 \) eV can also be calculated from our data.\(^{3} \) Note that both values are significantly stronger than \( D^0(VN\text{-H}) \) and much more similar to \( D^0(N\text{-H}) \) = 3.46 eV. This is consistent with the \( VN^+\text{-H} \) and \( VN^+\text{-NH} \) structures rather than \( H^+\text{-VN} \) and \( H^+\text{-VN} \). Also the value of \( D^0(VN^+\text{-H}) \) can be combined with IP(VN) and IP(H) = 13.598 eV to calculate the proton affinity of VN, PA(VN). Depending on the value of IP(VN) used, the result is either 9.66 ± 0.20 eV = 223 ± 5 kcal/mol or 9.79 ± 0.29 eV = 226 ± 7 kcal/mol. Both values are in good agreement with BGF’s direct measurement of 225 ± 6 kcal/mol.\(^{3} \) Actually, BGF determined that VN* did proton transfer to diethylamine but not to pyridine. Using updated values for the proton affinities of these molecules, 225.9 and 220.8 kcal/mol,\(^{38} \) respectively, it can be seen that our values for PA(VN) are in excellent agreement with the BGF experimental results.

Bond Energy–Bond Order Correlation. In a previous study,\(^{12} \) we have pointed out that a qualitative feeling for the bonding character in simple transition-metal-ion–ligand species can be obtained by a comparison of their bond strengths with analogous bonds in carbon systems. We found that there was a reasonable correlation between \( V^+\text{-CH} \) \( (n = 0-3) \) and \( CH_2^-\text{-CH}_2 \) bond strengths. Figure 7 shows this correlation for the updated thermochemistry given in Table IV and the present results for \( V^+\text{-NH}_n \) \( (n = 0-2) \) compared to both \( D^0(CH_2^-\text{-NH}_2) \) and \( D^0(CH_2^-\text{-NH}_2^-) \). The latter comparison is included to demonstrate that the

\[
\Delta H^o = D(M^+\text{-L}) - D(M^-\text{-L}) - (M^-\text{-L}) \quad (11)
\]

**TABLE IV: Heats of Formation and Bond Dissociation Energies (eV) at 298 K**

<table>
<thead>
<tr>
<th>M^+\text{-L}</th>
<th>ΔH^o(M^+\text{-L}) ( \text{eV} )</th>
<th>ΔD(M^+\text{-L}^-) ( \text{eV} )</th>
<th>ΔD(M^+\text{-L}-\text{L}^-) ( \text{eV} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>V^+\text{-H}</td>
<td>12.32 (0.10)</td>
<td>2.09 (0.06)</td>
<td>4.55 (0.02)</td>
</tr>
<tr>
<td>V^+\text{-CH}</td>
<td>13.30 (0.10)</td>
<td>5.00 (0.06)</td>
<td>9.95 (0.18)</td>
</tr>
<tr>
<td>V^+\text{-CH}_2</td>
<td>12.68 (0.17)</td>
<td>3.47 (0.14)</td>
<td>7.47 (0.06)</td>
</tr>
<tr>
<td>V^+\text{-CH}_3</td>
<td>11.49 (0.13)</td>
<td>2.17 (0.10)</td>
<td>3.89 (0.02)</td>
</tr>
<tr>
<td>V^+\text{-N}</td>
<td>12.40 (0.10)</td>
<td>4.65 (0.06)</td>
<td>9.66 (0.20)</td>
</tr>
<tr>
<td>V^+\text{-NH}</td>
<td>11.55 (0.18)</td>
<td>4.30 (0.16)</td>
<td>6.32 (0.16)</td>
</tr>
<tr>
<td>V^+\text{-NH}_2</td>
<td>10.93 (0.13)</td>
<td>3.10 (0.17)</td>
<td>5.19 (0.11)</td>
</tr>
<tr>
<td>V^+\text{-NH}_3</td>
<td>11.55 (0.18)</td>
<td>2.88 (0.19)</td>
<td>4.82 (0.02)</td>
</tr>
</tbody>
</table>

* Ion heats of formation are calculated by using the thermal electron convention. \( \Delta H^o(V) = 12.15 \) (0.08) eV, ref 15. * Values are from this study unless otherwise noted. Bond energies for these species are calculated by using heats of formation from ref 15 and: Podley, J.; Brey, J.; Fowler, R. D.; Kivel, S. P. Thermochernical Data of Organic Compounds. 2nd. ed.; Chapman and Hall: London, 1986. The only exception is that \( \Delta H^o(CH_2^-\text{-NH}_2) = 32 ± 2 \) kcal/mol was taken from ref 38. Reference 25. Reference 16. Reference 3.

Figure 7 shows the bond strength of \( V^+\text{-N} \) agrees readily with the trend seen for the carbon ligands and falls in the region associated with covalent triple bonds. Since \( V^+ \) has four valence electrons, a triple bond implies that VN* should have a single unpaired electron on the metal and thus should have a double bond ground state. Indeed, Kunze and Harrison calculate that the ground state is \( ^2 \)Δ containing a \( V^-\text{-N} \) triple bond.\(^{33} \)

In contrast, the bond strengths of \( V^+\text{-NH}_2 \) and \( V^+\text{-NH} \) are markedly higher than their isoelectronic analogues, \( V^+\text{-CH}_2 \) and \( V^+\text{-CH} \). Thus, the nitrogen lone pair electrons must be contributing to the bonding, which implies a significant overlap between the lone pair and the empty metal d orbitals. In effect, this appears to form an additional bond worth about 1.4 eV (based on the \( 1 / 2 \) intercept of the upper line in Figure 7). The end result is a \( V^+\text{-NH}_2 \) bond energy comparable in strength to a \( V^+\text{-C} \) double bond and a \( V^+\text{-NH} \) bond energy nearly equal to the \( V^+\text{-N} \) triple bond. (Using an identical analysis, BGF came to the same conclusion that VN* had only a double bond,\(^{16} \) in part because they plotted their lower limit of 4.0 eV on a plot similar to Figure 7.) This bonding picture implies that VN* and \( VN^+ \) have triplet and quartet ground states, respectively.

**Reaction Mechanism.** In this section, the information presented above is used to postulate a mechanism for the reaction of \( V^+ \) with
ammonia. This mechanism must not only explain the observed products but also the dependence of the reactions on electronic excitation. The mechanism we favor is directly analogous to that proposed to explain the interaction of \( V^+ \) with methane.\(^{16} \)

A simple reaction mechanism for the interaction of \( V^+ \) with ammonia involves oxidative addition of the N–H bond at the metal center to form intermediate I, H–V\(^+\)–NH\(_3\). Bond additivity arguments can be used to estimate that the reaction leading to this intermediate is \( \approx 0.6 \) eV exothermic. Intermediate I clearly explains the observed competition between the VH\(^+\) and VNH\(_2\)+ products. Formation of VNH\(_2\)+ dominates at low energies because it is thermodynamically favored. Formation of VH\(^+\) dominates at high energies since it is strongly favored by angular momentum constraints. This argument has been detailed previously\(^{16} \) and can be seen qualitatively by comparing the reduced masses of the reactants and products. For reaction 3, the reduced masses of the reactants, \( \mu(\text{VNH}_2^++\text{H}) = 1.0 \) amu, and the products, \( \mu(\text{VH}^++\text{NH}_3) = 12.2 \) amu, are nearly equal and it is therefore easy to conserve orbital angular momentum. However, for reaction 6, the reduced mass of the products, \( \mu(\text{VH}^++\text{NH}_3) = 1.0 \) amu, is much smaller such that orbital angular momentum cannot be conserved for large reactant impact parameters. The result is that the latter channel is more highly restricted than the former channel. Reactions 4 and 7 can be explained by decomposition reactions involving the products formed via intermediate I, as mentioned above.

Intermediate I is also an obvious first step in the dehydrogenation reaction. We envision two possible mechanisms for the dehydrogenation step starting from I: via intermediate II or the four-center transition state III. There are three reasons that we think II is an unlikely pathway for reaction 5. First, by using bond additivity arguments, we estimate that the production of II from \( V^+(\alpha^3\Sigma^+) + \text{NH}_3 \) is endothermic by \( \approx 0.2 \pm 0.2 \) eV. Since formation of VNH\(_2\)+ occurs exothermically with no activation barrier, this argument discounts intermediate II as a viable pathway (but not definitively). Secondly, BGF have observed that the analogue of reaction 5 also occurs for Sc\(^+\),\(^{44} \) and we have verified that the behavior of the Sc\(^+\) + ammonia system is very similar to that of \( V^+ \).\(^{30} \) Since Sc\(^+\) has only two valence electrons, it cannot support the three covalent bonds of II. Even if a the Sc\(^+\)-H bonds are covalent and the HSc\(^+\)-NH bond is dative, formation of the scandium analogue of II is estimated to require \( 0.6-2 \) eV in excess of the Sc\(^+\) + NH\(_3\) reactants. Third, using arguments similar to the first two, the H\(_2\)V\(^+\)-CH\(_4\) intermediate analogous to II has been discounted as an intermediate for dehydrogenation of methane (and here the first argument is much more definitive).\(^{16} \) Since the state-specific behavior of the reactions of \( V^+ \) with methane and ammonia parallel one another, it seems reasonable that the mechanisms are also analogous.

If we use the same three criteria to evaluate intermediate III, we conclude it is a viable pathway for dehydrogenation. First, bond additivity estimates suggest that formation of III requires no energy in excess of the reactants (a reverse activation barrier of \( 0.12 \pm 0.16 \) eV). While this may at first appear to be a small activation barrier for a four-center reaction, ordinarily symmetry forbidden, calculations have demonstrated that such an elimination can be facile if the transition-metal ligand bonds are covalent and have significant d character.\(^{40} \) Both of these conditions are probably met by intermediate I. Second, the analogous four-center intermediate in the scandium system will have comparable energetics to III since only two metal electrons are ever involved.

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\(^{16} \) Clemmer et al., J. Phys. Chem., submitted for publication.


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Electronic-State Dependence. In order to understand the effects of electronic states upon product formation, molecular orbital ideas can be employed. These arguments have been used successfully before in the reactions of atomic metal ions with \( \text{H}_2 \)\(^2\) and \( \text{CH}_4 \)\(^2\) elucidating reactivities and mechanisms. Oxidative addition of \( \text{N}–\text{H} \) to a metal center is achieved by donation of electrons in \( \sigma \) bonding orbitals into empty \( 4s \) and \( 3d \) \( \sigma \) orbitals of the metal and back-donation of the metal \( 3d \) electrons into \( \sigma^* \) antibonding orbitals. This increases the electron density between the metal and molecular fragments and also lengthens the \( \text{N}–\text{H} \) bond. For \( V^+ \), the states having \( 3d^4 \) configurations, e.g., \( \alpha^3\Pi \) and \( \alpha^3\Delta \), and higher triplet states, can interact with NH\(_3\) in this way and therefore should have attractive potential energy surfaces. However, if the metal \( 4s \) orbital (and to a lesser extent the \( 3d \) orbital) is occupied, the surfaces will be repulsive since the \( 4s \) orbital correlates to an antibonding orbital of the intermediate. Thus, states such as \( \alpha^3\Pi(4s3d^2) \) and \( \alpha^3\Delta(4s3d^2) \) are anticipated to be less reactive than those with empty \( 4s \) and \( 3d \) orbitals.

Further insight into the \( V^+ \) with ammonia system can be gained by examining the potential energy surfaces involved. In order to do this, the spin state of intermediate I must be established. If I truly contains covalent \( \text{V–H} \) and \( \text{V–N} \) bonds, then two of the four valence electrons on \( V^+ \) are involved in bonding. Therefore, the ground state of I must be a triplet. Based on these ideas, the calculations shown in Figure 8 can be drawn. This diagram shows an initial interaction for all states that is attractive due to the ion–dipole potential. The depth of this \( V^+–\text{NH}_3 \) well is estimated to be about the same as that for \( \text{Na}^+–\text{NH}_3, 1.3 \) eV,\(^{41} \) since these ions have similar ionic radii,\(^{42} \) although the well depth could be
somewhat larger.43 (In all likelihood, the interaction energy is smaller for V⁺ states having occupied 4s orbitals than for those which do not. In the absence of any definitive information on this effect, we use a uniform well depth for all states.) Both the a²F and the a'²F states have repulsive surfaces at closer distances since the 4s orbital is occupied. The a''²D surface is presumably less repulsive than these states since it has a 3d² configuration; but this state has the wrong spin to produce ground-state H₂VN⁺H². Therefore, this surface also rises in energy at close interaction distances. The a¹P state (which we let represent other higher lying states as well) has both the correct spin and electron configuration to smoothly generate higher lying states as well) has both the correct spin and electron configuration to smoothly generate higher lying states as well. This corresponds to moving the electron originally in the 4s orbital into a 3d orbital to remove the repulsive interactions.

Now consider the spin states of the dehydrogenation products, H₂(¹Σ⁺) + VN⁺H⁺, presumed to have a triplet ground state as noted above. Therefore, reaction of the quintet surfaces to form VN⁺H⁺ must occur via a crossing from a quintet surface to a triplet surface. Such a crossing exists, Figure 8, and presumably would require spin-orbit coupling of these surfaces. The enhanced reactivity of the triplet states is a result of favorable energy and spin, Figure 8. The relative inertness of the quintet states (~0.6% as reactive as the triplet states) suggests that such spin-orbit coupling is rather inefficient in this system. Similar results were found for the reaction of V⁺ with methane.15

For the endothermic production of VH⁺ and VN⁺H₂⁺, the ground states of the products are VH⁺ (1A¹g), VN⁺H₂⁺ (1Σ⁺g), and H(2S) + VN⁺H₂⁺, presumed to have a quartet ground state as noted above. Thus, these reactions are spin-allowed from both quintet and triplet states of V⁺. Because of this, VH⁺ and VN⁺H₂⁺ formation need not be as sensitive to the reactant state as the dehydrogenation process. However, these channels also show large magnitude increases for the triplet states (most specifically, the a²F) reactivity compared to the quintet states. This strong state dependence for VH⁺ and VN⁺H₂⁺ formation implies that the reaction proceeds predominantly through intermediate I. Note that, if the critical intermediate is the simple adduct, then the potential energy surfaces of Figure 8 would predict no strong spin dependence to these reactions. The intermediacy of I, however, easily explains this dependence.

Summary
The results of this study indicate that the most likely reaction mechanism for the V⁺ + NH₃ system proceeds via oxidative addition of a N-H bond to yield I, H⁺-V⁻-NH₂⁺. Simple bond cleavage forms VH⁺ and VN⁺H²⁺ in endothermic processes, while four-center molecular elimination of H₂ leads to the exothermic formation of VN⁺H++. At higher kinetic energies, VN⁺H²⁺ decomposes to form VN⁺H⁺ + H and VN⁺ + H₂.

The triplet states are found to be much more reactive than the quintet states even after accounting for differences in available energy. This is shown by the approximate state-specific cross sections shown in Figure 6. While there are sizable uncertainties in the determination of the absolute excited-state cross sections, the qualitative differences in reactivity are clear and unambiguous. This behavior is rationalized by using molecular orbital arguments which have been tremendously useful in understanding the reactions of atomic transition metals with H₂ and CH₄. These lead to the qualitative potential energy surfaces for the V⁺ + ammonia system shown in Figure 8. These surfaces show that the triplet states can more easily form the triplet ground state of the insertion intermediate I because of spin conservation and thermochemistry. Some reactions of the quintet states of V⁺ (certainly dehydrogenation, reaction 5, and any other processes evolving through the ground state of intermediate I) must involve spin-forbidden processes. These reactions presumably occur via spin-orbit mixing, which is apparently inefficient in this system as it was for the reaction of V⁺ with CH₄.

Thermochemistry for VH⁺ is found to agree with previous results. New thermochemistry for VN⁺, VN⁺H⁺, and VN⁺H₂⁺ is reported and tabulated in Table IV. Comparison of the ammonia system to analogous organic systems shows that the interaction of the lone pair electrons on the nitrogen atom with empty metal d orbitals leads to substantial increases in the bond strengths of the VN⁺H+ and VN⁺H₂+ species. Thus, these molecules have bond strengths which correspond most closely to a triple and a double bond, respectively. Also, the VN⁺ bond energy implies that this molecule is triply bound.

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Registry No. V⁺, 14782-33-3; NH₃, 7664-41-7.

Structure, Energetics, and Vibrational Spectrum of H₃N⁺HOH

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The geometry of H₃N⁺HOH is fully optimized by using a number of basis sets including 6-31G** and variants which add a second set of d-functions and a diffuse sp-shell. The internal geometries are altered very little as a result of complexation with the exception of a 0.008 Å stretch of the bridging N-H bond; N-H-O is within 5° of linearity. The electronic binding energy of the complex is 5.7 kcal/mol, 1.2 kcal/mol of which is a result of electron correlation. The calculations confirm the expected red shift and intensification of the ν₅ band of the proton donor molecule within the complex. All of the intermolecular modes are calculated and used to help assign bands in the experimental spectrum.

I. Introduction
Due in part to the weakness and polar character of the hydrogen bond, its properties are sensitive to perturbation by its environment. For this reason, recent examination of this interaction in the gas phase¹⁵ and in low-temperature inert gas matrices⁸⁻¹⁰ has revealed

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