Pulsed field-ionization photoelectron-photoion coincidence study of the process \( \text{N}_2 + h\nu \rightarrow \text{N}_2^+ + \text{N} + e^- \): Bond dissociation energies of \( \text{N}_2 \) and \( \text{N}_2^+ \)

Xiaonan Tang, Yu Hou, and C. Y. Ng

Department of Chemistry, University of California, Davis, California 95616

Branko Ruscic

Chemistry Division, Argonne National Laboratory, Argonne, Illinois 60639-4831

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We have examined the dissociative photoionization reaction \( \text{N}_2 + h\nu \rightarrow \text{N}_2^+ + \text{N} + e^- \) near its threshold using the pulsed field-ionization photoelectron-photoion coincidence (PFI-PEPICO) time-of-flight (TOF) method. By examining the kinetic-energy release based on the simulation of the \( \text{N}_2^+ \) PFI-PEPICO TOF peak profile as a function of vacuum ultraviolet photon energy and by analyzing the breakdown curves of \( \text{N}_2^+ \) and \( \text{N}_2 \), we have determined the 0-K threshold or appearance energy (AE) of this reaction to be 24.2884 ± 0.0010 eV. Using this 0-K AE, together with known ionization energies of \( \text{N}_2 \) and \( \text{N}_2^+ \), results in more precise values for the 0-K bond dissociation energies of \( \text{N}--\text{N}^+ \) (9.7543 ± 0.0010 eV) and \( \text{N}--\text{N}^+ (8.7076 ± 0.0010 \text{ eV}) \) and the 0-K heats of formation for \( \text{N}_2^+ \) (112.469 ± 0.012 kcal/mol) and \( \text{N}_2^+ \) (447.634 ± 0.012 kcal/mol). © 2005 American Institute of Physics. [DOI: 10.1063/1.1995699]

I. INTRODUCTION

One of the most valuable thermochemical data that can be obtained in single-photon vacuum ultraviolet (vuv) photoionization-photoelectron measurements is the 0-K threshold or appearance energy (AE) of a dissociative photoionization process. The recent introduction of the high-resolution pulsed field-ionization photoelectron (PFI-PE) photoion coincidence (PFI-PEPICO) time-of-flight (TOF) method has allowed the determination of 0-K AE values for many dissociative photoionization processes of simple molecules with unprecedented precision (=0.1 kJ/mol), which is limited mostly by the precision achieved in PFI-PE measurements.3–14

As pointed out in previous PFI-PEPICO TOF studies, the analysis of the breakdown curves measured in PEPICO TOF experiments represents one of the most reliable methods for 0-K AE determinations.1–4 Because the 0-K AE is distinctly marked by the disappearance energy of the parent ion observed in the breakdown diagram regardless the temperature of the gas sample, the 0-K AE value can be determined without any simulations.1–4 Furthermore, the PFI-PEPICO TOF peak for the daughter ion contains information about the kinetic-energy release.12 By measuring the kinetic-energy release based on the simulation of the daughter PFI-PEPICO TOF peak as a function of vuv photoionization energy or internal excitation of the parent ion, we have previously identified the dissociative photoionization for the formation of \( \text{CO}^+ + \text{O} \) and \( \text{CO}^+ + \text{O} \) from \( \text{CO}_2 \).12

Here, we report the result of a PFI-PEPICO TOF study of the dissociative photoionization process,

\[
\text{N}_2 + h\nu \rightarrow \text{N}_2^+ + e^- \rightarrow \text{N}_2^+ + \text{N} + e^-.
\]

When the excited diatomic cation such as \( \text{N}_2^+ \) with a fixed internal energy dissociates near the dissociation threshold such that the atomic fragments \( \text{N}^+ + \text{N} \) are produced in their ground states, there can only be one single kinetic-energy release because the atomic fragments cannot possess rovibrational energies. As a result, the analysis of the measured PFI-PEPICO TOF peak of the daughter \( \text{N}^+ \) ion by varying the vuv photoionization energy (\( h\nu \)) becomes a very sensitive mean for the determination of the 0-K AE of process (1). The combined kinetic-energy release and breakdown curve measurements have yielded a highly precise 0-K AE(\( \text{N}^+ \)) from \( \text{N}_2 \). Because the ionization energies (IEs) for \( \text{N} \) and \( \text{N}_2 \) have been very precisely measured,15–17 the 0-K bond dissociation energies (\( D_0 's \)) for \( \text{N}--\text{N} \) and \( \text{N}^+--\text{N}^- \) can be obtained by the relations,

\[
D_0 (\text{N}^+--\text{N}) = \text{AE}(\text{N}^+) - \text{IE}(\text{N}_2),
\]

\[
D_0 (\text{N}--\text{N}) = \text{AE}(\text{N}_2^+) - \text{IE}(\text{N}).
\]

The primary motivation for the present investigation is to try to establish an improved value for \( D_0 (\text{N}--\text{N}) \), and hence \( \Delta H_f (N) \). The current experimental value18–21 for \( \Delta H_f (N) \) is derived from \( D_0 (\text{N}--\text{N}) = 78.715 \pm 0.50 \text{ cm}^{-1} \) (225.057 ± 0.143 kcal/mol) (Refs. 22–25) obtained by Buttenbender and Herzberg. The need to improve this quantity has surfaced recently while building and analyzing the Core (Argonne) Thermochemical Network [C(A)TN], which is currently being developed as a separate effort and is the basis for Active Thermochemical Tables (ATcT).26,27 ATcT, are a new approach that allows the derivation of accurate, reliable, and internally consistent thermochemical values. In contrast with traditional (sequential) thermochemistry, ATcT extract
the thermochemical values of interest by statistically analyzing, iteratively optimizing, and solving a thermochemical network (TN) that is constructed from all the relevant thermochemical determinations and explicitly exposes and utilizes the interdependencies between various thermochemical quantities. Besides producing thermochemical values that are based on full and optimal utilization of the aggregate knowledge stored in the TN, ATcT offer a number of additional features that are neither present nor possible in the traditional approach, such as easy propagation of new knowledge with a concomitant automatic update of all affected thermochemical values, availability of the full covariance matrix detailing the interdependencies of values, a hypothesis testing and evaluation capability, etc. In particular, an inspection of the computed statistical measures (such as the covariance matrix and the sensitivity matrix) that are obtained by analyzing and manipulating the current C(A)TN, combined with insights into the TN topology, can lead to the isolation of “weak” links in the network, thereby generating pointers to the highly desired experimental or computational determinations that will efficiently improve the knowledge content of the TN.26,27 One such current “weak link” is the value of \[D_0(N-N),\] which is the primary contributor to the resulting value of the enthalpy of formation of atomic nitrogen, \(\Delta H_f^o(N)\) and presently limits its accuracy. The latter enthalpy of formation is one of the global “key” thermochemical values, and is the pivoting quantity for the establishment of accurate enthalpies of formation of NO and NO\(_2\) through \[D_0(\text{NO})\] and \[D_0(O-\text{NO}).\] Propagating further through the inherent thermochemical interdependencies, \(\Delta H_f^o(\text{NO})\) and \(\Delta H_f^o(\text{NO}_2)\) are in turn pivoting quantities responsible for the enthalpies of formation of the other nitrogen oxides, NO\(_x\), further affecting the thermochemistry of the chemical species with the formula \(H_xN_yO_z\), and so on. The value (and the uncertainty) for \(\Delta H_f^o(N)\), is also of pivotal importance as a fixed reference value for all electronic structure computations of that derive the enthalpy of formation through computed atomization energies (such as the high-accuracy composite methods from the G1 family developed by Curtiss et al.\(^{28}\) or the Wn family recently developed by Martin and de Oliveira\(^{29}\)). Clearly, in all these cases the value and uncertainty of \(\Delta H_f^o(N)\) affects the value and uncertainty of the enthalpies of formation of the target nitrogen-containing chemical species, and, in fact, this contribution scales linearly with the number of nitrogen atoms in the target species.

II. EXPERIMENTAL CONSIDERATIONS

The PFI-PEPICO TOF experiments were carried out using the high-resolution vuv photoelectron-photoion facility of the chemical dynamics beamline at the advanced light source\(^{30}\) (ALS) which was operated in the multibunch mode (synchrotron ring period=656 ns, dark gap=84 ns). The facility consists of a 10-cm period undulator, a gas harmonic filter, a windowless 6.65-m off-plane Eagle mounted monochromator, and a multipurpose photoelectron photoion apparatus.\(^{13,11,23}\) In the present experiment, helium was used in the harmonic gas filter to suppress higher undulator harmonics with photon energies greater than 24.59 eV. A 2400-lines/mm grating (dispersion=0.64 Å/mm) was used to disperse the first-order harmonic of the undulator vuv beam with entrance/exit slits set in the range of 20–100 µm. The resulting monochromatic vuv beam was then focused into the photoionization/photoexcitation (PI/PEX) region of the photoelectron-photoion apparatus.

The photon energy calibration was achieved using the Ne\(^{+}(2P_3/2, 2P_1/2)\) and He\(^{+}(3S)\) PFI-PE bands. The calibration scheme assumes that the Stark shifts of the ionization energies (IEs) of \(N_2\) and rare gases are identical. On the basis of previous experiments, the accuracy of the energy calibration is shown to be within ±0.5 meV.\(^{33}\)

The schematic diagram of photoelectron-photoion coincidence spectrometer, typical voltages applied to individual lenses, and the procedures for PFI-PEPICO measurements have been described in detail previously.\(^{2}\) The PFI-PEPICO TOF spectra were measured in the photon energy region 23.58–24.35 eV. A dc field of 0.2 V/cm was used at the PI/PEX region to disperse the prompt electrons. The PFI at the PI/PEX region was achieved by applying an electric pulse (height=9 V/cm, width=200 ns) at the dark gap of the synchrotron period. The PFI electric pulse also served to extract PFI photoions toward the ion detector.\(^{2}\) The average accumulation time for one PFI-PEPICO spectrum is about 50–60 min. The PFI-PEPICO resolution achieved is 1.0 meV [full width at half maximum (FWHM)]. The \(N_2\) sample was introduced into the PI/PEX region as a continuous, neat \(N_2\) supersonic beam at a total stagnation pressure of 760 Torr. The molecular beam was skimmed by a circular skimmer (diameter=1 mm) before entering the photoionization chamber. The alignment of the nozzle and the skimmer was adjusted to give the lowest translational temperature (=5 K) for \(N_2\) as measured by the width of the \(N_2^+\) PFI-PEPICO TOF peak. During the experiment, the beam source chamber and the photoionization chamber were maintained at \(5 \times 10^{-4}\) and \(5 \times 10^{-7}\) Torr, respectively.

III. RESULTS AND DISCUSSION

Two methods are adopted to determine the 0-K \(AE\) of \(N^+\) from \(N_2\) [process (1)]: One is based on the measured breakdown curves for \(N^+\) and \(N_2^+\), and the other relies on the kinetic energy release analysis of the observed PFI-PEPICO TOF peak profiles for \(N^+\) and \(N_2^+\).

A. Breakdown curves for \(N^+\) and \(N_2^+\)

An illustrative set of selected PFI-PEPICO TOF spectra obtained at \(h\nu=23.5810, 24.2672, 24.2871, 24.2881, 24.2888, \) and 24.2891 eV, are shown in Figs. 1(a)–1(f). At 23.5810 eV, which is well below the dissociative photoionization threshold of process (1), only the parent TOF peak is observed at 20.85 µs. The spectrum at 24.2672 eV [Fig. 1(b)] reveals the fragment \(N^+\) TOF peak at 14.65 µs in additional to the parent ion TOF peak. The broadness of this \(N^+\) TOF peak indicates that the fragment \(N^+\) ions result mostly from dissociative photoionization of ambient thermal \(N_2\) neutrals in the photoionization chamber. The exceptionally high intensity observed for this \(N^+\) TOF peak is consistent with the lifetime switching effect\(^{5}\) at energies very slightly
The process $N_2 + h\nu \rightarrow N^* + N + e^-$ since $N^* + H_2O \rightarrow$ thermal $N_+ signals are found to increase to 42% and 50% of pulse. Here $N^*$ and $N_2$ prior to PFI by the application of the PFI electric-field photoionization energies of observed of the PFI-PEPICO TOF peak for the fragment $N_+ in Fig. 1 be seen to consist of contributions from a broad thermal $N_2$ threshold for process well above the IE$N_2$, the PFI-PEPICO TOF peak for fragment $N_+$ can be seen to consist of contributions from a broad thermal $N_2$ component and a narrow supersonically cooled $N_2$ component and becomes dominant compared to that for $N_2^*$ [see Fig. 1(c)]. The PFI-PEPICO TOF spectra observed at vuv energies $\geq 24.2881$ eV reveal only a negligibly small TOF peak for $N_2^*$ as shown in Figs. 1(d)–1(f). A quantitative analysis of the TOF profile at 23.5810 eV shows that the thermal $N_2^*$ signal is about 10%–15% of the $N_2^*$ signal. The thermal $N^*$ signals are found to increase to 42% and 50% of the total $N^*$ and $N_2^*$ signal at 24.2672 and 24.2871 eV, respectively. The thermal $N^*$ signal drops to 23% and 15% of the total $N^*$ and $N_2^*$ signal at 24.2888 and 24.2891 eV, respectively.

The respective integrated ion signals under the PFI-PEPICO TOF peaks for $N^*$ and $N_2^*$ were taken as the intensities of $N^*[I(N^*)]$ and $N_2^*[I(N_2^*)]$. The fractional abundances of $N^*$ and $N_2^*$ are determined as $I(N^*)/[I(N^*) + I(N_2^*)]$ and $I(N_2^*)/[I(N^*) + I(N_2^*)]$, respectively. The breakdown curves for $N^*$ and $N_2^*$, i.e., the fractional abundances for $N^*$ (open circles) and $N_2^*$ (solid circles) plotted as a function of $h\nu$ in the range of 24.25–24.29 eV, are depicted in Fig. 2. As expected, the fractional abundance for the parent $N_2^*$ ion decreases as the $h\nu$ is increased with the concomitant increase in the fractional abundance of the daughter $N^*$ ion. We note that the breakdown curves for $N^*$ and $N_2^*$ constructed based on the intensities determined by the total TOF peak area have included contributions from both the cold and thermal components of $N_2$. The dashed lines are simulated breakdown curves obtained using a procedure described previously and assuming an effective rotational temperature of 300 K for $N_2$. Since the PFI-PEPICO TOF spectra resolve the dissociation due to cold $N_2$ from those of thermal $N_2$, we have also constructed the breakdown diagram based only on the intensities of the cold $N^*$ and $N_2^*$ components. The open and closed rectangular symbols in Fig. 2 show the respective fractional abundances of $N^*$ and $N_2^*$ determined by using only the cold components of $N^*$ and $N_2^*$. As expected, the breakdown curves constructed based on the dissociation of cold $N_2^*$ formed by photoionization of supersonically cooled $N_2$ are much sharper. The solid curves of Fig. 2 represent the simulated curves obtained by assuming a rotational temperature of 10 K for the cold component of $N_2$. As pointed out in previous PFI-PEPICO studies, the
0-K AE can be unambiguously determined by the disappearance energy of the parent ion. The dispersion of hot electrons into the dark gap of the synchrotron ring period can give rise to coincidence background for N$_2^+$, In such cases, we have shown that the disappearance energy is manifested as a sharp break in the breakdown curve for the parent ion, where the breakdown curve of the parent ion reaches its lowest value and levels off. The break in the breakdown curve for the parent N$_2^+$ ion is clearly observed at 24.288 eV, where the intensity of the TOF peak for parent N$_2^+$ is found to be within the noise level. We note that the dark gap (84 ns) of the synchrotron ring period used in the present experiment is narrower than that (104 ns) used previously. Consequently, the small residual N$_2^+$ signal (fractional abundance $\approx 0.03$) observed at energies $\approx 24.288$ eV is likely due to the dispersion of prompt electrons into the dark gap of the synchrotron ring period. Taking into account the error of $\pm 0.5$ meV for vuv energy calibration, we conclude from this set of measurements that the 0-K AE(N$_2^+$) for process (1) is 24.2888±0.0010 eV, where the uncertainty reflects an estimated 95% confidence interval.

In PFI-PEPICO measurements of polyatomic systems, where the density of states is high, the lifetime switching effect is manifested as a step in the PFI-PE spectrum at the 0-K AE of the dissociative photoionization process. In the study of CO$_2$, a peak in the PFI-PE spectrum is found to coincide with each of the 0-K AE for the formation of O$^+(4S) +$CO($1\Sigma^+$) and CO$^+(2\Sigma^+) +$O($3P$) from CO$_2$. Similarly, peaks are observed in the PFI-PE spectrum at energies coinciding with the 0-K AE for the dissociative photoionization processes CS$_2 +$hv $\rightarrow$ S$^+(3S) + $CS($1\Sigma^+$; $v = 0 \text{ and } 1)$ + e$^-$. We have measured the PFI-PE spectrum for N$_2$ in the region of 23.40–24.40 eV using a PFI-PE resolution of 5 cm$^{-1}$ (FWHM). The PFI-PE spectrum of N$_2$ thus observed reveals many vibrational bands of the N$_2^+$ (X, C, and 2 $^2\Pi_0^+$) states. These PFI-PE vibrational bands are found to superimpose on a continuous component of PFI-PE signals with significant intensities, indicating that the PFI-PE spectrum is mediated by dissociative states of N$_2$. Figure 3 depicts the PFI-PE spectrum of N$_2$ recorded in the region of 24.20–24.40 eV. The prominent PFI-PE peak at 24.338 eV can be ascribed to the vibrational band of the N$_2^+(C)$ state. Other minor peaks could have contributions from vibrational bands of the N$_2^+(2^2\Pi_u)$.

### B. Kinetic-energy release analysis

When a dc field $E$ (V/cm) is maintained at the photoionization region, the width ($\Delta T$) of the N$^+$ PFI-PEPICO TOF peak can be calculated as

$$\Delta T = \frac{\sqrt{8m(E_{KE} + E_T)}}{qe},$$

where, $m$ and $q$ represent the mass and the electric charge of N$^+$, respectively; $E_{KE}$ is the kinetic energy gained by N$^+$ due to the energy release of reaction (1); and $E_T$ represents the thermal translational energy of N$^+$, which is determined by the translational temperature of N$_2$ along the TOF axis. Equation (4) can also be expressed as

$$E_{KE} = \frac{q^2E^2}{8m}(\Delta T)^2 - E_T.$$

In the present experiment, instead of using a dc electric field, a pulsed electric field was employed to the repeller to extract the N$^+$ and N$_2^+$ ions. The kinetic energy for N$^+$ can be obtained by simulation of the observed TOF peak using the SIMION program. The simulation indicates that Eq. (4) can be used to calculate the kinetic energy for N$^+$ with only minor discrepancies if $E$ is replaced by the effective dc field ($E_{eff}$) of 2.74 V/cm, which is calculated as (amplitude of the PFI pulse $\times$ PFI pulse width $100$ ns)/synchrotron ring period of 656 ns.

The parent N$_2^+$ ion in PEPICO measurements are all produced in the PFI of N$_2^+(m)$ formed by vuv excitation of N$_2$. Thus, the cold component of the N$_2^+$ PFI-PEPICO TOF peak is expected to be decided solely by the N$_2$ supersonic beam temperature, which does not change with respect to $h\nu$. This expectation is supported by the observation that the FWHM for the PFI-PEPICO TOF peak of N$_2^+$ in the $h\nu$ range from 24.257 to 24.287 eV is constant (see Fig. 4). As indicated above, the TOF peaks for N$_2^+$ are dominated by contributions from N$_2$ in the supersonic beam. The simulation$^{12}$ of the TOF peak profiles for parent N$_2^+$ ions observed in the vuv energy range of 24.257–24.287 eV yields a translational temperature of $\approx 5$ K for the supersonically cooled N$_2$ beam.

The FWHMs observed for the TOF peaks of N$^+$ in the vuv energy range of 24.254–24.2977 eV are also depicted in Fig. 4. The high FWHM values for the N$^+$ TOF peaks observed at 24.257–24.287 eV are consistent with the fact that the N$^+$ ions produced at these energies are contributed predominantly by the dissociative photoionization of thermal N$_2$ existing in the photoionization chamber. As the $h\nu$ is increased, the dissociation from the supersonically cooled N$_2$ component increases, resulting in narrower FWHMs for TOF peaks of N$^+$. The TOF peak width should reach its minimum value at the 0-K AE of process (1). The further increase in...
produce an excited N\(_2\) involves two consequential steps: H\(_2\)O\(_8\)/H\(_2\)O\(_9\) the disappearance energy of N\(_2\), and the observed total kinetic-energy release from N\(_2\) translational temperature of 5 K for N\(_2\). The process N\(_2^+\)→N\(_+^+\)+N\(_++\) ion and a zero kinetic-energy electron) and 1(b) (N\(_2^+\) dissociation to produce N\(_+^+\)+N), the appearance energy and the observed total kinetic-energy release (KER=2E\(_{\text{KE}}\)) for process (1) at energy h\(_v\) are related as

\[ \text{AE}(N^+) = h\(_v\) - 2E_{\text{KE}}. \]  

(6)

The magnified views of the PFI-PEPICO TOF peaks observed for N\(_+^+\) at h\(_v=24.2883, 24.2888, 24.2907, 24.2938, \) and 24.3365 eV are shown in Figs. 5(a)–5(e), respectively. We have simulated\(^{12}\) these TOF peaks for N\(_+^+\) (produced from N\(_2^+\) at a translational temperature of 5 K) to determine the kinetic-energy release (E\(_{\text{KE}}\)). The best-fitted curves (solid line) are also shown in Figs. 5(a)–5(e). The KER values determined at h\(_v=24.2883, 24.2888, 24.2907, 24.2938, \) and 24.3365 eV are given in Table I. The 0-K AE values calculated using Eq. (6) at individual h\(_v\) values are also listed in Table I. The mean 0-K AE is 24.2880 eV with a standard deviation of the sample of 0.0008, the standard deviation of the mean of 0.0003, and the 95% confidence limit interval of 0.0009 eV.

C. Thermochemistry

On the basis of the breakdown curves for N\(_+^+\) and N\(_2^+\) or the disappearance energy of N\(_2\) observed in the PFI-PEPICO

![FIG. 4. Comparison of the full width at half maximum (FWHMs) of the N\(_+^+\) (filled circles) and N\(_2^+\) (open circles) PFI-PEPICO TOF peaks in the range of 24.257–24.300 eV. The FWHMs of the N\(_2^+\) TOF peaks correspond to a translational temperature of 5 K for N\(_2\).](image)

![FIG. 5. Selected PFI-PEPICO TOF spectra for N\(_+^+\) at the VUV photon energies of (a) 24.2883, (b) 24.2888, (c) 24.2907, (d) 24.2938, and (e) 24.3365 eV. The simulated spectra are shown as solid lines. The N\(_+^+\) peaks are centered at 14.65 µs.](image)

TOF measurements, the 0-K AE(N\(_+^+\)) from N\(_2\) is found to be 24.2888±0.0010 eV. By taking the narrowest width of the PFI-PEPICO TOF peak for N\(_+^+\) to mark the threshold for process (1), we obtain the 0-K AE(N\(_+^+\)) =24.2883±0.0010 eV. The KER analysis for process (1) as summarized in Table I gives the AE(N\(_+^+\)) =24.2880±0.0009 eV. From these, we recommended the final value of 24.2884±0.0010 eV for the 0-K AE(N\(_+^+\)) of process (1).

### TABLE I. Total kinetic-energy releases (KER) and 0-K AE values for reaction (1) measured at selected VUV photoionization energies (h\(_v\)).

<table>
<thead>
<tr>
<th>h(_v) (eV)</th>
<th>KER (eV)</th>
<th>0-K AE(^a) (eV)</th>
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<tr>
<td>24.2883</td>
<td>0.0003</td>
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<tr>
<td>24.2888</td>
<td>0.0013</td>
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<td>0.0035</td>
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</tr>
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<td>24.2938</td>
<td>0.0047</td>
<td>24.2891</td>
</tr>
<tr>
<td>24.3365</td>
<td>0.0481</td>
<td>24.2884</td>
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</table>

\(^a\)Calculated using Eq. (6).
\(^b\)Average value for the 0-K AE; the quoted uncertainty is the corresponding 95% confidence interval.
Using the latter 0-K AE(N⁺) and the known IE(N₂) =125 668.00±0.25 cm⁻¹ or 15.580 85±0.000 03 eV (Ref. 17) and IE(N)=14.534 14 eV (Ref. 15), one straightforwardly obtains by sequential propagation \( D_0(N-N) = 8.7076±0.0010 \) eV and \( D_0(N-N)=9.7543±0.0010 \) eV. The latter bond dissociation energy in turn leads to 
\[ \Delta H_f(0)=112.469±0.012 \text{ kcal/mol} \] 
and, with IE(N), to 
\[ \Delta H_f(0)=447.634±0.012 \text{ kcal/mol}. \]

The optical excitation of \( N_2 \) in the \( ^1Σ_g^+ \) ground electronic state was found to lead to dissociation with a threshold at 12.139 eV. Thus, the \( D_0(N-N) \) can be calculated if the states of the product N atoms are known. Herzberg had narrowed down the possible product states to be \( N^+(2D) + N(2^3S) + N(2^3P) \), 16 which would yield the corresponding \( D_0(N-N) \) values of 9.756 and 7.373 eV. Assuming that the predissociation of the \( N_2(3^1Π_u) \) state produces \( N^+(2S) + N(2D) + N(2^3P) \), a value of 9.759 eV (9.757 eV) can be deduced. 16 The value \( D_0(N-N) = 9.7543±0.0010 \) eV obtained in the present study, while in relatively good agreement (i.e., within their uncertainty) with \( D_0(N-N)=78.715±50 \text{ cm}^{-1} \) or 9.754 08±0.000 03 eV reported 12,12 by Buttenbender and Herzberg, represents an important experimental refinement of this quantity. The currently accepted values for \( \Delta H_f(N) \) and \( \Delta H_f(N⁺) \) (e.g., those found in the JANAF Thermochemical Tables 20 or in the NIST WebBook 5) are based on the \( D_0(N-N) \) value of Buttenbender and Herzberg, as originally suggested by the CODATA Review 19 of Cox et al., which recommends \( \Delta H_f(298.15) = 112.970±0.096 \text{ kcal/mol} \), implying \( \Delta H_f(0) = 112.528±0.096 \text{ kcal/mol} \). Our new value, \( \Delta H_f(0) = 112.469±0.012 \text{ kcal/mol} \), obtained by sequential propagation of the measured appearance energy of \( N^+ \) from \( N_2 \), is slightly lower and about eight times more accurate than the currently used value. Inserting the present measurements of the appearance energy of \( N^+ \) from \( N_2 \) into the C(A)TN, together with other relevant determinations, and analyzing and solving the full complement of thermochemical interdependencies using Active Thermochemical Tables leads to a very similar value for \( \Delta H_f(N) \). This fully optimized recommended value obtained through ATcT (and the related discussion) will be the topic of a separate forthcoming report. 40

IV. SUMMARY

We have conducted a high-resolution PFI-PEPICO TOF study of the dissociative photoionization process of \( N_2 \). The 0-K AE(N⁺) values for process (1) deduced based on the breakdown curves of N⁺ and \( N_2^+ \) and the KER analysis of the N⁺ PFI-PEPICO TOF peaks are in agreement. We recommend the average value of 24.2884±0.0010 eV for the 0-K AE(N⁺) from \( N_2 \). The latter value has allowed the determination of more precise values for the \( D_0(N-N), D_0(N^+N), \) \( \Delta H_f(0), \) and \( \Delta H_f(0) \). (Argonne). One of the authors (C.Y.N.) also acknowledges partial supports by the AFOSR Grant No. F49620-03-1-0116 and NSF ATM 0317422. Another author (B.R.) also acknowledges partial support and effort from the numerous team members of the Collaboratory for Multi-Scale Chemical Science (CMCS), which is a project within the National Collaboratories Program sponsored by the U.S. Department of Energy’s Division of Mathematical, Information, and Computational Sciences. Portions of this research are related to the effort of a Task Group of the International Union of Pure and Applied Chemistry (2003-024-1-100), which focuses on the thermochemistry of chemical species implicated in combustion and atmospheric chemistry.

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The process \( \text{N}_2 + h^+ \rightarrow \text{N}^+ + \text{N} + e^- \) 

<table>
<thead>
<tr>
<th>Reference</th>
<th>Description</th>
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