Electron-Induced Radiolysis of Astrochemically Relevant Ammonia Ices

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Abstract
We elucidate mechanisms of electron-induced radiolysis in cosmic (interstellar, planetary, and cometary) ice analogs of ammonia (NH₃), likely the most abundant nitrogen-containing compound in the interstellar medium (ISM). Astrochemical processes were simulated under ultrahigh vacuum conditions by high-energy (1 keV) and low-energy (7 eV) electron-irradiation of nanoscale thin films of ammonia deposited on cryogenically cooled metal substrates. Irradiated films were analyzed by temperature-programmed desorption (TPD). Experiments with ammonia isotopologues provide convincing evidence for the electron-induced formation of hydrazine (N₂H₄) and diazene (N₂H₂) from condensed NH₃. To understand the dynamics of ammonia radiolysis, the dependence of hydrazine and diazene yields on incident electron energy, electron flux, electron fluence, film thickness, and ice temperature were investigated. Radiolysis yield measurements versus (1) irradiation time and (2) film thickness are semi-quantitatively consistent with a reaction mechanism that involves a bimolecular step for the formation of hydrazine and diazene from the dimerization of amidogen (NH₂) and imine (NH) radicals, respectively. The apparent decrease in radiolysis yield of hydrazine and diazene with decreasing electron flux at constant fluence may be due to the competing desorption of these radicals at 90 K under low incident electron flux conditions. The production of hydrazine at electron energies as low as 7 eV and an ice temperature of 22 K is consistent with condensed phase radiolysis being mediated by low-energy secondary electrons produced by the interaction of high-energy radiation with matter. These results provide a basis from which we can begin to understand the mechanisms by which ammonia can form more complex species in cosmic ices.

Key words: radiation chemistry, interstellar ice, cosmic rays, photochemistry, NH₂ radicals, NH radicals

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1 Introduction

1.1 Astrochemical relevance and motivation for investigating ammonia

Ammonia (NH$_3$) is of astrochemical interest due to its presence in diverse extraterrestrial environments. For example, the microwave sounding measurements made in August 2016 by the Juno spacecraft provided convincing evidence for an ammonia-rich plume in the atmosphere of Jupiter.$^1$ Significant amounts of ammonia have been detected in the atmospheres of Saturn,$^2$ Uranus,$^3$ and Neptune,$^4$ as well as on the surface of Saturn’s moon, Enceladus.$^5$ Additionally, cometary ices in the Halley, Borrelly, Hartley-Good, and Thiele comets exhibit substantial ammonia abundances (> 10%) relative to water.$^6$ Studies of interstellar ice composition in protostellar regions indicate that with the possible exception of molecular nitrogen, that ammonia is the most prevalent nitrogen-containing compound in interstellar ices, with an abundance of 1 – 10% relative to water.$^7$ Gas phase NH$_2$ radicals discovered in the interstellar medium$^8$ may result from radiation processing of ammonia ices or from ground state N atoms reacting with ammonia in the solid phase.$^9$ Moreover, the relatively high abundance of ammonia in star-forming regions and elsewhere suggests that it may be a precursor for N-containing prebiotic molecules. The discovery in 2016 of glycine in the coma around Comet 67P/Churyumov-Gerasimenko by Rosetta’s ROSINA spectrometer is significant since comets are thought to have preserved icy grains initially present in the interstellar medium prior to star formation.$^{10}$ Cyanomethanimine (NC$_2$HNH), a C–N bond containing prebiotic molecule recently discovered in an interstellar gas cloud (Sgr B2(N)),$^{11}$ is of particular interest because experimental and theoretical studies suggest that it cannot be formed in the gas phase.$^{12}$ Therefore, understanding the energetic processing of ammonia ices may provide insights into how complex prebiotic
species form in condensed phase environments within the interstellar medium. While our recent work characterizes the photochemistry of condensed ammonia,\textsuperscript{13} this work focuses on the other major form of energetic processing: radiation chemistry.

1.2 Radiation chemistry of cosmic ices

While low-temperature (\(~40\) K) thermal processing of organic ice mixtures to form more complex species has been demonstrated,\textsuperscript{14} photolysis/radiolysis of extraterrestrial ices is thought to be the primary path to prebiotic molecules in cosmic ices. Except for rare entities such as Titan which has an icy surface consisting of methane and ethane, the dominant component of extraterrestrial ices is water. Celestial ices, some containing organics, have been detected on solar-system bodies such as asteroids (e.g., 24 Themis), comets (e.g., 67P/Churyumov-Gerasimenko), planets (e.g., Neptune), dwarf planets (e.g., Ceres), and moons (e.g., Europa). In addition, cosmic ices also include interstellar ices within dark, dense molecular clouds. These interstellar ices, having a thickness of a few hundred molecular layers, are composed of primarily of water (H\textsubscript{2}O), carbon dioxide (CO\textsubscript{2}), methanol (CH\textsubscript{3}OH), and ammonia (NH\textsubscript{3}) surrounding carbonaceous or silicaceous micron-size dust particles.\textsuperscript{15} Far (Deep)-UV (200 – 300 nm), near-UV (300 – 400 nm), and visible (400 – 700 nm) light are incident upon these cosmic ices and can initiate photochemistry. While photochemistry proceeds via electronic excitation by absorbed photons, radiation chemistry, by definition, involves ionization. In addition to condensed phase photochemistry, radiation chemistry in the condensed phase occurs in outer space because extraterrestrial ices are also continuously bombarded by high-energy particles (e.g., cosmic rays consisting mostly of protons) and high-energy photons (e.g., vacuum-UV (100 – 200 nm), extreme-UV (10 – 100 nm), X-rays, and \(\gamma\)-rays). Given that the ionization energy of a generic molecule is \(~10\) eV and that the ionization threshold is reduced in the condensed phase
compared to the gas phase, Lyman alpha radiation (10.2 eV) may result in ionization and subsequent radiation-induced chemistry. Results of numerous studies suggest that condensed phase radiolysis is mediated by low-energy (< 20 eV) electrons produced by the interaction of high-energy radiation with matter. The role of low-energy electrons in the radiolysis of cosmic ices has been recently reviewed. Interestingly, in 2018, glycine formation was observed in CO₂:CH₄:NH₃ ices irradiated by sub-ionization electrons.

While in the sections below we focus on radiolysis of condensed ammonia, we note that several studies have identified hydrazine but not diazene as a radiolysis product of gaseous ammonia.

1.3 High-Energy Ion Irradiation of Condensed Ammonia

The high-energy (~500 MeV) heavy ion (⁶⁴Ni²⁺ and ⁷⁰Zn²⁺) bombardment of pure ammonia ices has been studied to mimic some cosmic ray-induced processes. Molecular hydrogen (H₂), molecular nitrogen (N₂), cis-diazone (HNNH), iso-diazone (NNH₂), hydrazine (N₂H₄), azide ion (N₃⁻), ammonium ion (NH₄⁺), and ammonium azide salt (NH₄⁺N₃⁻) were all observed following high-energy ion irradiation of condensed pure ammonia (molecular structures shown in Figure 1). The sole radical reported was the amidogen (amido) radical (NH₂). Radiolysis by 0.8 MeV protons of ammonia/water ices and by 144 keV S⁹⁺ ions of ammonia/carbon dioxide ices have also been studied.

1.4 High-Energy Photon Irradiation of Condensed Ammonia

Following irradiation of NH₃ ice films by 150 eV photons, results of Near-Edge X-ray Absorption Fine Structure (NEXAFS) spectroscopy confirmed the production of large amounts of N₂ (12%), as well as cis-diazone (HNNH), iso-diazone (NNH₂), azide (N₃⁻), and hydrazoic
acid (hydrogen azide) (HN₃).²⁷ Hydrazine was not detected in this study likely due to limitations associated with NEXAFS.²⁷

1.5 High-Energy Electron Irradiation of Condensed Ammonia

The radiolysis of condensed ammonia, initiated by high-energy electrons, has been studied using post-irradiation temperature programmed desorption (TPD) experiments.²⁸,²⁹,³⁰⁻³¹ Results of such experiments conducted following 5 keV electron irradiation of ammonia ice films indicate the production of molecular nitrogen (N₂), molecular hydrogen (H₂), hydrazine (N₂H₄), and diazene (N₂H₂).²⁸ Based on the observation of peaks in m/z = 32 (N₂H₄⁺), m/z = 31 (N₂H₃⁺), m/z = 30 (N₂H₂⁺), and m/z = 29 (N₂H⁺) at ~ 160 K in post-irradiation TPD experiments, hydrazine was identified as a product of high-energy electron irradiation of condensed ammonia.²⁸ The presence of peaks in m/z = 30 and 29 at temperatures above 160 K, even after the m/z = 32 and 31 signals had decreased to zero, was attributed to diazene.²⁸ Hydrazoic acid (HN₃) was not detected in post-irradiation temperature programmed desorption experiments likely because it is thermally unstable, and decomposes at a lower temperature than its desorption temperature.²⁸ Results of recent post-irradiation TPD experiments, employing reflectron time-of-flight mass spectrometry, indicate the production of triazane (N₃H₅) and cyclotriazane/triazene (N₃H₃) following 5 keV electron irradiation of condensed ammonia.³⁰⁻³¹

In addition to the aforementioned TPD measurements, infrared spectroscopy was used to verify the 5 keV electron-induced production of both the cis- and iso- isomers of diazene from condensed ammonia.²⁸ Trans-diazene formation could not be confirmed, probably because the vibrational features are too similar to those of ammonia.²⁸ Two additional products, hydrazoic acid (HN₃) and the amidogen radical (NH₂), were identified through infrared spectroscopy in the same study.²⁸ It is believed that hydrazine is not detectable in infrared studies as its signature
bands overlap with those exhibited by ammonia, with the possible exception of a feature at 895 cm\(^{-1}\).\(^{24,28}\)

In this article, we describe the dynamics of hydrazine and diazene production from the radiolysis of condensed ammonia by high-energy (1 keV) and low-energy (7–10 eV) electrons. First, we confirm the identification of these two radiolysis products by using three isotopologues (NH\(_3\), ND\(_3\), and \(^{15}\)NH\(_3\)) of ammonia. Second, we demonstrate that electron-induced reactions likely occur in the multilayer and not on the metal surface based on qualitatively similar results obtained on both Mo(110) and Ta(110). Third, we provide experimental evidence consistent with two-step mechanisms that involve bimolecular steps for the electron-induced synthesis of hydrazine and diazene from condensed ammonia. Finally, we establish that hydrazine results from condensed phase reactions of ammonia initiated by electrons with incident energies as low as 7 eV, suggesting a role for dissociative electron attachment or electron impact excitation, or both. This observation is qualitatively consistent with the now widely accepted view that low-energy electrons play a pivotal role in high-energy radiolysis.

2 Experimental
Experiments at Wellesley College were conducted in a custom-designed stainless steel ultrahigh vacuum (UHV) chamber, described in detail previously.\(^{32}\) The base pressure of the UHV chamber is maintained at \(\sim 1 \times 10^{-9}\) Torr and is monitored with a nude ion gauge. In order to maintain ultrahigh vacuum pressures, a turbomolecular pump, rotary vane mechanical pump, ion pump, and titanium sublimation pump are employed. An electrically isolated single crystal (Ta(110) or Mo(110)) substrate inside the chamber is cooled via a liquid nitrogen reservoir to cryogenic temperatures of \(\sim 90\) K. Surface temperature was monitored using a W/5\% Re vs. W/26\% Re thermocouple spot welded to the edge of the crystal. The crystal is fixed to a
manipulator that is capable of $\theta$, $x$, $y$, and $z$ movement. Typically, experiments necessitated only rotation between the gas doser, electron gun, and mass spectrometer.

Samples of NH$_3$ (99.99%, Matheson), ND$_3$ (99 atom% D, Cambridge Isotope Laboratories), and $^{15}$NH$_3$ (98 atom % $^{15}$N, Aldrich) were used without further purification. Films of were prepared by introducing gas samples first into a “manifold” or “dosing chamber,” where pressure was measured by Baratron (MKS Instruments), and then via a precision leak valve, into the UHV chamber to condense upon the Mo(110)/Ta(110) substrate. Sample film thickness, which for convenience is expressed in monolayers (ML) of ammonia, was varied by controlling the change in pressure within the dosing chamber during film deposition. The quantity of gas (i.e., pressure change) required to form 1 ML was determined from temperature programed desorption measurements with unirradiated ammonia films. 1 ML was defined to be the maximum exposure of ammonia that did not yield a multilayer peak. This method to determine film thickness has been used in other laboratories studying cosmic ice analogs.$^{33-34}$ Note, however, that the ML designation relates more properly to the quantity or mass of deposited NH$_3$, rather than to the thickness of a film, which can vary with porosity and film order, both of which are dependent on substrate temperature and other parameters of film deposition.

Films were irradiated using a Kimball Physics FRA2X1-2 flood electron gun, at energies between 5 eV and 1000 eV and transmitted electron currents varying between 1.5 $\mu$A and 35 $\mu$A, corresponding to electron fluxes of $1.9 \times 10^{13}$ electrons cm$^{-2}$s$^{-1}$ and $4.4 \times 10^{14}$ electrons cm$^{-2}$s$^{-1}$. The energy distribution of the electron gun has a FWHM of $\sim 0.4$ eV. The entire 1 cm$^2$ crystal was irradiated during electron irradiation.

The Mo(110) surface was cleaned by radiative heating using radiation emitted by a hot tungsten filament positioned immediately behind the metal crystal. Further heating by electron
bombardment to ~2200 K ensured the desorption of remaining nitrogen and hydrogen between experiments. Once daily, the surface was further cleaned of carbon by dosing oxygen (5 \times 10^{-9} Torr for five minutes) at ~1200 K and subsequently heating to ~2200 K to remove excess adsorbed oxygen. The Ta(110) substrate was cleaned by radiative and electron bombardment heating to 2200 K following each experiment.

Post-irradiation TPD experiments were conducted by radiatively heating the crystal surface from ~90 K to ~800 K while monitoring three to five mass spectral fragments of the desorbing species with a triple-filtered Hiden IDP Series 500 quadrupole mass spectrometer. TPD experiments conducted in the absence of electron irradiation served as control experiments.

Because surface temperatures below 90 K were not attainable at Wellesley College, additional experiments were conducted at the University of Sherbrooke under UHV conditions (at a base pressure ~6\times10^{-11} Torr) in a previously described experimental system. Target films were formed by vapor deposition onto a Pt-foil substrate, the temperature of which can be controlled between 22 and 400 K with a closed-cycle He cryostat and resistive heating elements. The thickness of films in monolayers (ML) condensed onto the substrate was determined with an absolute accuracy of ~30% by a volumetric dosing procedure similar to that described for the Wellesley experiments. A chromel-alumel thermocouple spot welded to the Pt foil facilitated the accurate measurement of film temperatures for TPD. The foil was cleaned between experiments by resistive heating to ~1000 K. For the experiments described in this work, NH₃ (99.9% MSD isotopes) was deposited on the Pt-foil substrate cooled to 22 K. Sample films were irradiated isothermally at 22 K with electrons of specific energies from an electron gun (Kimball Physics ELG-2). The energy resolution of this electron beam is approximately 0.5
eV. For post-irradiation TPD measurements, the entire film was irradiated in raster-mode (1.25 \times 10^{12} \text{ electrons cm}^{-2} \text{ s}^{-1} \text{ flux onto a sample of 1 cm}^2).

For TPD measurements, desorbing species were ionized and detected with a quadrupole mass spectrometer (Extrel 150-QC). Samples on the Pt foil were heated resistively by currents of up to 7 A delivered by a floatable power supply (Kepco ATE 25-10M). The latter was controlled by an in-house LabVIEW program to provide a typical linear heating rate of 10 K min\(^{-1}\). The mass spectrometer ionizer is off during electron-beam irradiation and requires ~ 30 minutes to stabilize once it is turned on. During this period, samples (i.e., both irradiated and unirradiated films) are rotated away from the mass spectrometer and are biased at −87 V relative to ground, to repel electrons from the ionizer of the mass spectrometer. This sample bias is maintained once samples are returned into position and throughout the heating process.

3 Results and Discussion

3.1 Identification of Hydrazine and Diazene

Hydrazine (N\(_2\)H\(_4\)) identification was based on results of TPD experiments conducted after 1000 eV electron irradiation of condensed ammonia (NH\(_3\)), showing desorption peaks at ~155 K for \(m/z = 32 (\text{N}_2\text{H}_4^+)\), 30 (\text{N}_2\text{H}_2^+), and 29 (\text{N}_2\text{H}^+) \text{ (Figure 1)}.\(^b\) The peak associated with \(m/z = 32 (\text{N}_2\text{H}_4^+)\) is the largest, consistent with this fragment being the dominant ion in the published hydrazine mass spectrum.\(^{38}\) The identification of hydrazine was further verified by isotopic labeling experiments whose results show the expected shift to \(m/z = 36 (\text{N}_2\text{D}_4^+), 34 (\text{N}_2\text{D}_3^+), \text{ and } 32 (\text{N}_2\text{D}_2^+) \text{ in irradiated deuterated ammonia (ND}_3\) \text{ (Supporting Information: Figure B: post-irradiation TPD of ND}_3\) \text{ as well as } m/z = 34 (^{15}\text{N}_2\text{H}_4^+), 32 (^{15}\text{N}_2\text{H}_2^+), \text{ and } 31 (^{15}\text{N}_2\text{H}^+) \text{ in }

\(^b\) We attribute to ion-molecule reactions of ammonia in the mass spectrometer the observed peaks at ~115 K for all monitored mass spectral fragments. These peaks appear because of the significant amount of ammonia present in our films. The ion-molecule reactions of ammonia are also detected in unirradiated films (data shown in Supporting Information, Figure A: TPD of unirradiated ND\(_3\) films).
irradiated $^{15}\text{NH}_3$ (Supporting Information: Figure C: post-irradiation TPD of $^{15}\text{NH}_3$). As discussed previously, prior studies have also identified hydrazine as a high-energy radiolysis (~550 MeV energy ions $^{24}$ and 5 keV electrons $^{28}$) product of ammonia.

![Figure 1](image)

**Figure 1**

Results showing production of hydrazine and diazene in temperature-programmed desorption experiments conducted after high-energy electron irradiation of a 500 ML film of condensed NH$_3$ on a Mo(110) surface. The peaks at ~115 K attributed to ammonia are due to ion-molecule reactions in the ionizer of the mass spectrometer, as verified by separate temperature programmed desorption experiments in which $m/z = 17$ was monitored.

Diazene (N$_2$H$_2$) identification was based on the results of the same experiment, showing desorption peaks at ~185 K for $m/z = 30$ (N$_2$H$_2^+$) and 29 (N$_2$H$^+$) (Figure 1). Because the parent ion of diazene is $m/z = 30$ (N$_2$H$_2^+$), the lack of desorption peaks at ~185 K for 32 (N$_2$H$_4^+$) provides additional support for our identification of diazene. Results of post-irradiation experiments involving isotopically labeled ammonia were used to verify our identification of diazene. We observe the expected mass spectral fragment shifts to $m/z = 36$ (N$_2$D$_4^+$), $m/z = 34$ (N$_2$D$_3^+$) and 32 (N$_2$D$_2^+$) in irradiated ND$_3$ (Supporting Information: Figure B: post-irradiation...
TPD of ND₃) and to \( m/z = 31 (^{15}\text{N}_2\text{H}^+) \) and 32 \( (^{15}\text{N}_2\text{H}_2^+) \) in irradiated \(^{15}\text{NH}_3\) (Supporting Information: Figure C: post-irradiation TPD of \(^{15}\text{NH}_3\)). Diazene \((\text{N}_2\text{H}_2)\) has been identified previously as a high-energy radiolysis product of condensed ammonia.²⁴,²⁷-²⁸

To investigate the role of the metal substrate surface, ammonia radiolysis experiments were conducted with both Mo(110) and Ta(110) crystals. Both hydrazine and diazene were detected during TPD experiments following electron-induced radiolysis of ammonia films regardless of the substrate. Furthermore, the TPD peaks for both hydrazine and diazene appear qualitatively similar for Mo(110), Ta(110), and polished polycrystalline silver,²⁸,³⁰-³¹ suggesting that the substrate surface plays a minimal, if any, role in the results reported in this study.

### 3.2 Proposed Radiolysis Mechanisms for Hydrazine and Diazene Formation

From the results of high-energy electron and photon irradiation of condensed ammonia, several mechanisms have been suggested for the formation of particular radiolysis products. Most notably, hydrazine is expected from the dimerization of two amidogen radicals \((\text{NH}_2)\).²⁷-²⁸

\[
\text{NH}_2 + \text{NH}_2 \rightarrow \text{N}_2\text{H}_4
\]  

(1)

As described previously, following irradiation of condensed ammonia by 5 keV electrons, the amidogen radical has been identified by post-irradiation infrared spectroscopy.²⁸ As described in detail below, (1) dissociative electron attachment, (2) electronic excitation, and (3) dissociative ionization likely play a role in the synthesis of amidogen radicals during condensed phase ammonia radiolysis.

The radiolytic production of amidogen radicals from ammonia may proceed via *dissociative electron attachment* resonances at incident electron energies of \(~ 6\) and \(~ 10\) eV in the production \(\text{H}^-/\text{D}^-\):
\[ \text{NH}_3 + e^- \rightarrow \text{H}^- + \text{NH}_2 \]  \hspace{1cm} (2)

The hydride anion (H\(^-\)) is the dominant product of the interactions of low-energy electrons with gaseous ammonia.\(^{39-40}\) In the electron stimulated desorption of deuterated ammonia condensed on a platinum substrate, low-energy electron irradiation primarily yielded D\(^-\) anions in a process also attributed to dissociative electron attachment.\(^{41}\) The production of H\(^-\) ions during high-energy (150 eV) electron stimulated desorption from one-monolayer thick ammonia films may also involve dissociative electron attachment via low-energy secondary electrons produced by 150 eV electrons interacting with the metal surface.\(^{42}\) Because gas and condensed phase ammonia demonstrate resonances in the production/desorption of the H\(^-\)/D\(^-\) ion, concomitant resonances should exist in the electron-induced production of the complementary amidogen radical (NH\(_2\)/ND\(_2\)).

During radiolysis, in addition to dissociative electron attachment, we hypothesize that low-energy secondary electrons produced within bulk ammonia ice may induce \textit{electronic excitation}, leading to the production of ground state amidogen radicals:

\[ \text{NH}_3 \xrightarrow{e^-} \text{NH}_2(\tilde{X} \ ^2\text{B}_1) + \text{H} \]  \hspace{1cm} (3)

The above electron-induced excitation process is a viable mechanism for the production of amidogen radicals given the energy threshold of 4.6 eV for the analogous gas-phase spin-allowed photon-induced excitation process:\(^{43}\)

\[ \text{NH}_3 + h\nu \rightarrow \text{NH}_2(\tilde{X} \ ^2\text{B}_1) + \text{H} \]  \hspace{1cm} (4)
Following electron impact excitation, another available molecular dissociation process is dipolar dissociation, the process by which the resultant excited electronic state induces ion-pair formation.

The electron-induced dissociative ionization of ammonia may also produce NH$_2$ radicals. Results of electron stimulated desorption studies indicate the formation of H$^+$ as the primary positive ion formed during high-energy (1.5 keV) electron irradiation of ten-monolayer (ML) thick ammonia films adsorbed on an argon spacer layer:

\[ \text{NH}_3 + e^- \rightarrow \text{NH}_2^+ + 2e^- \]  \hspace{1cm} (5)

\[ \text{NH}_2^+ \rightarrow \text{H}^+ + \text{NH}_2 \]  \hspace{1cm} (6)

As shown above, the formation of H$^+$ from ionized ammonia involves the concomitant production of NH$_2$ radicals. Proton transfer (acid-base chemistry) is another route for the synthesis of NH$_2$ radicals via dissociative ionization of ammonia:

\[ \text{NH}_3^+ + \text{NH}_2 \rightarrow \text{NH}_2 + \text{NH}_4^+ \]  \hspace{1cm} (7)

Based on ultraviolet photoelectron spectra which show a 1.2 eV energy shift upon ammonia condensation, we estimate that the above dissociative ionization mechanism has an electron energy threshold of \(~8.9\) eV, which is less than the gas phase ionization energy of 10.1 eV for ammonia.

Given the copious numbers of low-energy (< 20 eV) electrons produced during high-energy radiolysis, low-energy secondary electron-induced processes (dissociative electron attachment, electron impact excitation, and electron impact ionization) likely play a critical role in the
condensed phase radiolysis of ammonia to form hydrazine via the dimerization of two amidogen radicals.

In contrast to hydrazine formation, according to a recent study involving 5 keV irradiation of condensed ammonia, diazene formation via the dimerization of two imine (NH) (imidogen) radicals was considered unlikely because the photon-induced gas phase formation of the NH radical ground $X^3\Sigma^-$ state from ammonia is spin-forbidden.28

$$\text{NH}_3 + h\nu \rightarrow \text{NH} (X^3\Sigma^-) + \text{H}_2 \quad (8)$$

Therefore, high-energy electron-induced diazene formation from condensed ammonia was attributed to the excitation of the hydrazine product to an unstable electronic state, followed by the removal of two hydrogen atoms:28

$$\text{NH}_2 + \text{NH}_2 \rightarrow \left[\text{N}_2\text{H}_4\right]^* \quad (9)$$

$$\left[\text{N}_2\text{H}_4\right]^* \rightarrow \text{N}_2\text{H}_2 + \text{H}_2 \quad (10)$$

An additional mechanism was proposed for the high-energy photon-induced formation of diazene from condensed ammonia. This radiolysis mechanism involves the N$_2$H$_3$ intermediate either reacting with itself or with an amidogen radical.27

Here, we propose, however, that the electron-induced condensed-phase production of diazene from the dimerization of two NH radicals is possible even though the photon-induced formation of the NH radical ground $X^3\Sigma^-$ state from gas phase ammonia is spin-forbidden. For example, the photon-induced formation of the excited NH ($a^1\Delta$) from gaseous ammonia is spin-allowed with a threshold photon energy of 5.7 eV.43 The photochemistry of gaseous ammonia may even produce excited imine radicals (e.g., NH ($A^3\Pi$)) via spin-forbidden transitions.43
More importantly, in general, electron-molecule interactions are not limited by spin conservation, as shown explicitly for ammonia. Because the electron-induced formation from ammonia of ground state and excited state imine radicals is likely, we suggest that radiolysis of condensed ammonia likely includes diazene formation via the dimerization of two imine (NH) radicals.

Although we are unable to make a distinction between yields formed by direct electron impact during irradiation and products formed during the heating stage after irradiation, it must be noted that diffusion of heavy radicals has been found at temperatures as low as ~ 35 K. Therefore, we expect that products observed in TPD experiments were formed during electron irradiation at 90 K.

In the sections below, we provide experimental evidence that is consistent with the dimerization of NH₂ and NH radicals producing N₂H₄ and N₂H₂, respectively. Specifically, we investigate the dependence of radiolysis yield on film characteristics (thickness and temperature) and irradiation parameters (electron energy, fluence, and flux).

### 3.3 Radiolysis yield as a function of film thickness

Results of post-irradiation TPD experiments conducted following irradiation at constant electron energy, electron flux, and electron fluence indicate that the yields of both hydrazine and diazene increase monotonically with increasing ammonia (ND₃) film thickness. To quantify the yield of hydrazine (N₂D₄), the m/z = 36 peak was integrated and the area under the curve was plotted against film thickness (Figure 3). The thinnest ammonia film in which hydrazine was detected was 15 monolayers. Because the first five data points show a semiquantitative quadratic relationship between hydrazine yield and the number of irradiated NH₃ molecules, this data is suggestive of a bimolecular reaction mechanism for the production of hydrazine. We attribute the
lack of quantitative agreement between the model and the experimental data to three factors: (1) difficulty in quantifying the role of the metal surface in quenching radiolytic processes in thin (< 25 ML) films; \(^{48}\) (2) the possible desorption of NH\(_2\) radicals during irradiation from 90 K ammonia ices that are thin (see below); and (3) the finite electron range, the straight-line penetration distance, in condensed ammonia causes the hydrazine yield to begin to saturate as the film thickness is increased beyond 25 monolayers. While the electron range is not available for solid NH\(_3\), many solids exhibit electron ranges for 1 keV electrons of between 20 and 100 nm (or approximately between 50 and 250 ML). \(^{49}\) The finite electron range in liquid water is calculated to be on the order of 10 nm. \(^{50}\)

![Figure 2](image)

**Figure 2**
Hydrazine (N\(_2\)D\(_4\)) yield monitored as a function of ammonia (ND\(_3\)) film thickness on Ta(110) surface. The first five data points were fit to a quadratic equation (red).

The quantification of diazene required additional calculations to ensure that the integration of the \(m/z = 32\) curve did not include any contribution from hydrazine (Figure 3).
This separation was accomplished by first multiplying the $m/z = 36$ curve by a factor such that this modified curve and the $m/z = 32$ curve would match for the first desorption peak. Finally, the area under the modified $m/z = 36$ curve was subtracted from the area under the $m/z = 32$ curve in order to quantify the area that was exclusive to the diazene product.

![Figure 3](image)

**Figure 3**
Calculating diazene yield from post-irradiation TPD data. The $m/e = 36$ mass spectra curve is multiplied by some factor in order to overlay its hydrazine peak at 124 K with its fragment peak in $m/e = 32$. The area under the $m/e = 36$ curve is subtracted from the area under the $m/e = 32$ curve. The gray color region shown corresponds to the hydrazine product. The white region between the black and red curves corresponds to the diazene product.

The yield of diazene as a function of film thickness (Figure 4) is also suggestive of a reaction mechanism involving a bimolecular step for the formation of diazene from two NH radicals.
Diazene (N$_2$D$_4$) yield monitored as a function of ammonia (ND$_3$) film thickness on Ta(110) surface. The first five data points were fit to a quadratic equation (red).

**Figure 4**

Electron Energy: 1000 eV
Flux: 2.1×$10^{12}$ cm$^{-2}$s$^{-1}$
Fluence: 2.1×$10^{14}$ cm$^{-2}$
3.4 Radiolysis yield as a function of fluence

Hydrazine (blue) and diazene (red) yields as a function of fluence. The yields of hydrazine and diazene were obtained by integrating areas under the $m/z = 36$ and 32 peaks, respectively. The $m/z = 32$ peak area was corrected as shown in Figure 3. ND$_3$ films (500 monolayers) were irradiated with 1000 eV electrons. Irradiation times were varied from 1 second to 60 seconds while the incident current was kept constant. The error bars represent standard deviations of the mean. The curves represent fits to a model described in the text.

The yield of hydrazine and diazene was monitored as a function of fluence (Figure 5) at constant film thickness by maintaining a constant electron flux ($4.1 \times 10^{13}$ electrons cm$^{-2}$s$^{-1}$) while varying the irradiation times. The yield functions demonstrate a high degree of similarity for irradiation times shorter than 20 sec ($8.1 \times 10^{14}$ electrons cm$^{-2}$). We attempted to fit the data using a two-step kinetic model, involving a bimolecular radical–radical dimerization between amidogen/imine radicals, as the formation mechanism for hydrazine/diazene:
In the above reaction mechanism, R represents the reactant ammonia, I is an intermediate species (NH$_2$ or NH), and P is the product (either hydrazine or diazene). Differential equations can be written for each step of this proposed mechanism:

\[
\frac{d[R]}{dt} = -k_1[R]
\]

\[
\frac{d[I]}{dt} = k_1[R] - k_2[I]^2
\]

\[
\frac{d[P]}{dt} = \frac{k_2[I]^2}{2}
\]

In the above equations, \(k_1 = F\sigma\), where \(F\) is the electron flux (electrons cm$^{-2}$s$^{-1}$) and \(\sigma\) is the total effective electron-induced degradation cross section. These coupled differential equations were solved numerically to generate the fits for the yields of hydrazine and diazene vs. electron fluence (Figure 5). We attribute the lack of quantitative agreement between the model and the experimental data to two factors: (1) the loss of product from the film during irradiation and (2) heating of the film prior to product quantification. Nevertheless, we suggest that the semi-quantitative agreement between our experimental data and the kinetic model is consistent with the two-step dimerization reaction mechanism proposed for the electron-induced formation of hydrazine and diazene from condensed ammonia. Although one must be cautious about over-interpreting parameters gleaned from such kinetic solid-state models involving coupled differential equations, we suggest a diazene formation model that is simpler than what has been
previously proposed.\textsuperscript{28} The dependence of the diazene yield on fluence is not consistent with an insertion reaction:

\begin{equation}
\text{NH} + \text{NH}_3 \rightarrow \text{N}_2\text{H}_4
\end{equation}

3.5 Radiolysis yield as a function of incident electron energy

Interestingly, hydrazine formation has been recently observed following 15 eV electron irradiation of condensed ammonia.\textsuperscript{51} Because our ultimate goal is to determine the role of low-energy electrons in the radiolysis of ammonia under astrochemically relevant conditions, we conducted TPD experiments following irradiation at lower incident electron energies. The lowest incident electron energy at which hydrazine and diazene production were consistently observed was 10 eV (Figure 6) at an ice temperature of 90 K, the lowest possible surface temperature attainable in the ultrahigh vacuum chamber at Wellesley College. These experiments involved four hours of electron irradiation. Even though a much higher fluence was used for the 10 eV experiments, the yield of hydrazine and diazene was considerably smaller than that for the 1000 eV electrons.
Results of post-irradiation TPD experiments conducted at the University of Sherbrooke provide evidence for the formation of hydrazine from ammonia ices at 22 K at incident electron energies as low as 7 eV (Figure 7). The production of hydrazine from condensed ammonia at sub-ionization electron energies suggests a role for one or both of electron impact excitation or dissociative electron attachment. These results are qualitatively compatible with the widely-accepted hypothesis that condensed phase radiolysis is mediated by low-energy electrons produced by the interaction of high-energy radiation with matter.
Figure 7
Post-irradiation TPD data following the irradiation of 20 monolayer NH$_3$ films with electrons with the indicated energies. Note that for measurements at 7 and 10 eV, the electron fluence was $\sim 2.5 \times 10^{18}$ electrons cm$^{-2}$ at a flux of $\sim 6.9 \times 10^{14}$ electrons cm$^{-2}$ s$^{-1}$. At higher energies the fluence was $\sim 3.8 \times 10^{18}$ cm$^{-2}$ at a flux of $\sim 1.1 \times 10^{15}$ electrons cm$^{-2}$ s$^{-1}$.

3.6 Radiation yield as a function of ice temperature

The influence of the ice temperature on ammonia radiolysis yield was investigated via TPD experiments conducted following 15 eV electron irradiation of ammonia ices at 22 and 77 K (Figure 8). This reduction in hydrazine yield with increasing ice temperature could be related to an increased desorption rate for NH$_2$ intermediates at temperatures approaching that of ammonia sublimation. Our attempt to measure electron stimulated NH$_2$ desorption at low and high
temperatures were hampered because the elevated partial pressure of NH$_3$ and small variations in film temperature around $\sim$77 K led to a large and constantly varying background signal for $m/z$=16. The reduction in hydrazine yield with increasing temperature could also be related to decreased dimerization of NH$_2$ species because of temperature-related changes in film morphology, porosity, and order. For example, the yields of D$_2$, O$_2$ and D$_2$O$_2$ in electron irradiated films of D$_2$O were found to be higher from porous amorphous samples, than from dense, crystalline ones$^{52}$ It was suggested that such behavior might be the result of lower recombination rates of dissociated molecules due to diffusion into pores, or due to the presence of L-defects in amorphous ice that may favor the formation of D$_2$O$_2$. The effects of film order and porosity thus merit further attention in a future study. Because higher temperatures likely lead to lower rate of N$_2$H$_4$ production and lower electron energies yield fewer radicals at equivalent fluxes, this radiolysis yield dependence on ice temperature likely contributes to the absence of detectable radiolysis products following electron irradiation of 90 K ammonia ices at an incident electron energy of 7 eV (data not shown). The longer electron range$^c$ of higher energy electrons$^{53}$ enhances the production of radical intermediates below the ice surface, facilitating the dimerization of NH$_2$ and NH intermediates to form hydrazine and diazene, respectively.

$^c$ Electron range, the straight-line penetration distance of electrons in matter, is typically calculated by using the continuous slowing down approximation (CSDA). Reference 50 shows that the electron range can increase by two or three orders of magnitude when the electron energy is increased from 10 eV to 1000 eV.
Figure 8
Post-irradiation TPD data following the irradiation of 25 monolayer NH$_3$ film with 15 eV electrons at two different ice temperatures, 22 K and 77 K.

Results of separate photochemistry studies of condensed ammonia support our hypothesis that NH$_2$ radicals desorb at an ice temperature of 90 K.$^{13}$ Hydrazine synthesis is observed at an ice temperature of 90 K following irradiation of ammonia ices with $<$ 7.4 eV photons which we estimate have a large mean free path of 0.2 microns, allowing for the synthesis of NH$_2$ radicals significantly below the ammonia ice surface, therefore reducing radical desorption.

3.7 Radiation yield as a function of electron flux

The effect of electron flux (dose rate) on the ammonia radiolysis product yield was investigated by conducting post-irradiation TPD experiments while keeping constant the electron energy (1000 eV), film thickness (500 ML), and ice temperature (90 K); the electron current and irradiation time were varied, but the total electron fluence (dose) was kept constant ($8.2 \times 10^{14}$
electrons cm$^{-2}$). Results of these experiments appear to show a dose-rate effect for the production of both hydrazine and diazene (Figure 9). The red curves are simply used to guide the reader’s eye and have no physical/mathematical basis. Below an incident current of $\sim 2.5 \mu A$, the yields of both hydrazine and diazene more or less increase with increasing incident flux (current) at constant fluence (total incident charge). This dependence on flux (dose rate) is surprising given that electrons are categorized as low linear energy transfer (LET) radiation for which the radiolysis yield depends only on radiation dose (fluence) and not on dose rate (flux).\textsuperscript{d} We attribute this unexpected result obtained at an ice temperature of 90 K to NH$_2$ and NH radical desorption which competes with the dimerization that yield hydrazine and diazene, respectively. At higher electron flux, radicals are more likely to be produced closer to each other, increasing the probability for dimerization while the probability for radical desorption remains unchanged. The yield vs. dose-rate curves eventually plateau likely because at higher fluxes the likelihood increases that the products are destroyed by subsequently arriving electrons.

\textsuperscript{d} In contrast, high LET radiation such as \( \alpha \) particles produce closely spaced ionizations which lead to a dose-rate effect.
Figure 9

ND$_3$ (500 monolayer) films were irradiated with 1000 eV electrons. Flux and irradiation times were varied during these experiments to ensure a constant fluence (dose). The yields of hydrazine (A) and the yields of diazene (B) are plotted as a function of flux (dose rate) at constant electron fluence (dose). Data was fit to a logarithmic function for visual guidance (red). The error bars represent standard deviations of the mean.

3.8 Astrochemical Implications

While gas-phase reactions and surface reactions on bare carbonaceous or silicaceous dust grains contribute to cosmic chemistry, energetic processing of cosmic ices via photochemistry and radiation chemistry is thought to be the dominant mechanism for the extraterrestrial synthesis of prebiotic molecules.$^{54}$ Discerning the role of photochemistry vs. radiation chemistry in astrochemistry requires a molecular-level mechanistic understanding these two processes. While the results discussed above provide insight into the radiolytic mechanisms for production of hydrazine (N$_2$H$_4$) and diazene (N$_2$H$_2$) from condensed ammonia, our preliminary TPD results (Supporting Information, Figure D: post-irradiation TPD of ND$_3$ showing) also show evidence for the production of triazane (N$_3$H$_5$) and cyclotriazane/triazene (N$_3$H$_3$) following high-energy (1000 eV) electron irradiation of condensed ammonia, consistent with previous studies.$^{30-31}$ The same reaction products are also observed following energetic processing of condensed ammonia by non-ionizing radiation consisting of < 7.4 eV photons.$^{13}$ This similarity, surprising given the
myriad differences between photon- and electron- induced processes,\textsuperscript{13} suggests that electronic excitation followed by dissociation to form radicals is the dominant mechanism for the photochemistry and radiation chemistry of condensed ammonia. The results presented in this publication are consistent with dimerization of radicals to form N-2 species following radiolysis of condensed ammonia. Studies such as these are necessary for incorporation into astrochemical models which since 2018 have begun to take into account condensed phase radiolysis induced by cosmic rays, allowing gas-phase chemistry, diffusive grain-surface chemistry, and ice-mantle photolysis and radiolysis to be simulated in tandem using rate equation-based modeling methods.\textsuperscript{55-56}

4 Conclusion

We provide a comprehensive report on condensed ammonia radiolysis as a function of (a) initial electron characteristics (energy, flux, and fluence) and (b) film features (thickness and temperature). Results of post-irradiation TPD experiments demonstrate the electron-induced production of hydrazine (N$_2$H$_4$) and diazene (N$_2$H$_2$) from ammonia condensed on both Mo(110) and Ta(110). Identification of these two radiolysis products was verified by conducting experiments with isotopologues of NH$_3$. Based on theoretical considerations, we argue that the electron-induced formation of NH radicals from condensed ammonia is not subject to spin conservation rules. From yield versus (1) irradiation time and (2) film thickness results, we suggest that both hydrazine and diazene are formed via two-step reaction mechanisms involving bimolecular steps for the dimerization of NH$_2$ and NH radicals, respectively. The enhanced desorption of these radical intermediates at an ice temperature of 90 K during electron irradiation contributes to (1) the surprising absence of detectable radiolysis products at incident electron energies below 10 eV and (2) the unanticipated dependence of the radiolysis yield on electron
dose rate at 90 K. At an ice temperature of 22 K, however, hydrazine formation occurs at incident electron energies as low as 7 eV, consistent with the widely-accepted hypothesis that low-energy secondary electrons mediate high-energy radiolysis in condensed matter.

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**Supporting Information**

1. Representative temperature-programmed desorption experiment of ND$_3$ performed in the absence of prior irradiation
2. Post-irradiation temperature-programmed desorption experiment of deuterated ammonia (ND$_3$)
3. Post-irradiation temperature-programmed desorption experiment of $^{15}$NH$_3$.
4. Evidence for the production of triazane (N$_3$H$_5$) and cyclotriazane/triazene (N$_3$H$_3$) following high-energy (1000 eV) electron irradiation of condensed ammonia.

**References**


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