國立中央大學

Physics Department
Master thesis

VUV and EUV irradiation of CH₄+NH₃ ice mixtures

- An implication of Tholin on Charon

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November 2017







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VUV and EUV irradiation of $\mathrm{CH_4}{+}\mathrm{NH_3}$ ice mixtures - An implication of Tholin on Charon

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VUV and EUV irradiation of $\mathrm{CH_4} + \mathrm{NH_3}$ ice mixtures - An implication of Tholin on Charon

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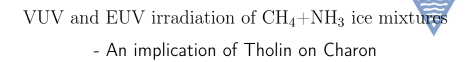
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- 欣逢中央大學教務處註冊組組長,蕭嘉璋老師,見微知著,並予協助,僅此誌謝。
- 承蒙太空及遙測研究中心蔡富安老師協助在Ubuntu 12.04上 TeXLive-2009測試成功,僅此誌謝。
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1. Methods

1.1 Laboratory Astrophysics

To study the chemical reactivity in astrophysical environment experimentally, we conducted our experiments in Interstellar photoprocessing system (IPS) [?], an ultrahigh vacuum chamber with base pressure 3×10^{-10} torr and 14 K, corresponds to a density of 10^6 cm⁻³, similar to dense cloud interiors. The system will be introduced in detail in section 1.1.1. To simulate the irradiation in interstellar environments, we use a micro-wave discharge hydrogen lamp (MDHL) and monochromatic extreme-ultraviolet irradiation (EUV) 30.4 nm to irradiate our ice mixtures, and they will be introduced in section 1.1.2 and 1.1.3 respectively. The experimental protocols will be elaborated in section 1.2. In order to better understand the physics behind, some basic theories of Infrared spectroscopy and concepts of chemical kinetics used in data analysis are included in section 1.3 and 1.4 respectively.

1.1.1 Experimental simulations by IPS system

We conducted our astrophysical simulations in Interstellar Photo Processing System (IPS) (figure 1.1). IPS consists in three systems: the main chamber, where our experiments take places; the detection system, where we collect our data; and a gasline system, where we prepare our samples.

The main system consists of an ultrahigh vacuum chamber equipped with a closed-cycle helium cryostat (CTI-M350). It is pumped by a turbo molecular pump (KTKT FF - 160/620ZE, capacity 600 liters s⁻¹), which is backed up by a scroll pump, and a non –evaporation getter pump. The getter pump is a powerful tool to adsorb residue gases inside the main chamber, with a larger surface area, H_2 , CO and N_2 are



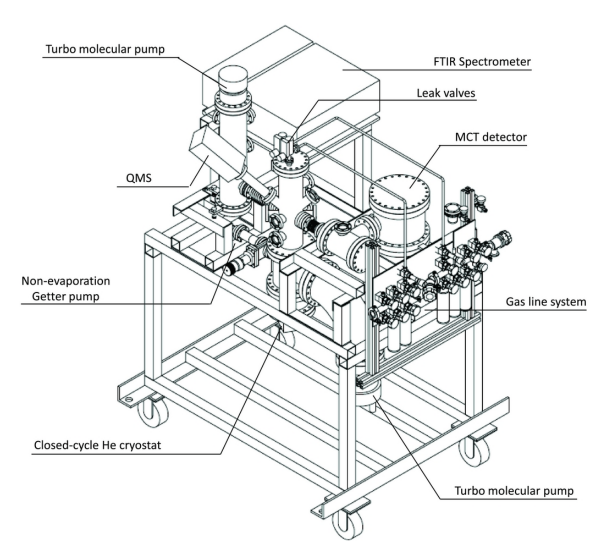


Figure 1.1: The schematic diagram of IPS system, mechanical pumps are not shown for clarity. (Quoted from Chen et al. 2014)

adsorbed to obtain a better base pressure. After baking, the base pressure of our main chamber can reach 1×10^{-10} torr at 14 K, monitored by a Granville-Phillips 370 Stabil-Ion gauge. This pressure can be used to demonstrate the dense cloud interior environments and star forming region. The substrate we have chosen is KBr, which can allow infra-red photons with 700 to 4000 cm^{-1} to penetrate. It is mounted by substrate holder made of oxygen-free copper, on the second stage of cold finger mounted on the tip of cryostat. Two silicon diodes and also a heater were placed onto the cold finger and one of the silicon diodes is near the substrate holder. They were connected to a temperature controller and PID system to achieve a warmup rate of 1K/min with an accuracy of 0.1 K.

The detection system consists in a mid-infrared Fourier transform spectrometer (mid-FTIR) (ABB FTLA2000-104) and a Quadrupole Mass Spectrometer (QMS). To prevent absorption bands of CO, CO₂ and H₂O gas in the atmosphere, the IR beam path was built inside vacuum, pumped by dry pump. The main chamber and the IR path are separated by ZnSe windows, which can allow infra-red penetration from 0.5 - 20 um with absorption less than 0.07 \%. In this study, the infrared spectra are obtained with resolution of $4~\mathrm{cm^{-1}}$ and averaged over 32scans. The angle between the IR beam path and the substrate holder is 45 degrees. The QMS (MKS Microvision 2) consists of a controller and mechanical part sealed by a mounting flange in ultrahigh vacuum. It is mounted 10 cm from the substrate and run with a resolution 0.5 a.m.u. The Ionizer release 70 eV electron by filament and ionize incoming molecules to positive charged ions between anode grid and repeller. The ions were accelerated by focus plate and enters ion filter, which consists of four circular rods, with a combination of A.C and D.C. potential to sieve whole bandpass ions at millisecond timescale. The selected ions enter ion detector and are detected by either faraday cup and continuous dynode electron multiplier (CDEM) which can secondary multiply weak signals.

The samples are prepared in situ in our gasline system. It contains four stainless steel bottles with the same volume, which is used to deter-



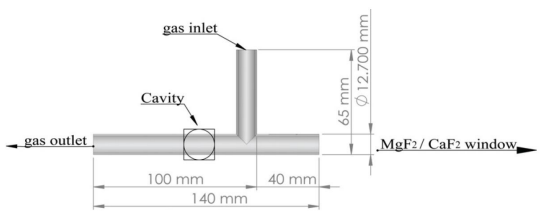


Figure 1.2: The cross-section of MDHL (T-type geometry) (Quoted from Chen et al. 2014).

mine relative proportion of the gas mixtures by their partial pressures. The ammonia gas 99.99% and methane 99.999% are mixed with partial pressure measured by a Baratron with 0 - 100 torr range with a 0.25

1.1.2 Vacuum-UV source

In order to simulate the photoprocessing of vacuum ultraviolet (VUV) irradiation onto the interstellar ices and ices on planetary bodies, including KBOs, the ice mixtures are irradiated with a T-type Microwave-Discharged Hydrogen-flow lamp (MDHL). The molecular hydrogen with pressure 0.4 torr flows through the lamp with a support of a mechanical pump. Using a 2.4 GHz microwave generator and high voltage discharge, a low pressure plasma is produced in the Evenson cavity. Figure 1.2 shows a cross-section of T-type quartz tube; the middle part of the T-type quartz tube is being tunned by a ceramic rod that is called Evenson cavity. In order to measure the photon flux in situ, we use an 88 % transmittance nickel mesh with its photoelectric efficiency being obtained by high-flux beamline in National Synchrotron and a SXUV 100 photodiode calibrated by NIST. A MgF₂ window is placed between the lamp and the sample holder to prevent penetration of VUV photons with wavelength shorter than 114nm, leads to a cut off at 114nm. Figure 1.3 shows a VUV emission spectrum of a MDHL. It consists in Ly- α (121.6nm) and H₂ molecular emission in 110-180 nm range. Chen et al. (2014) showed that the spectral characteristics of the VUV light emitted in this range depends on the gas type (mixture of

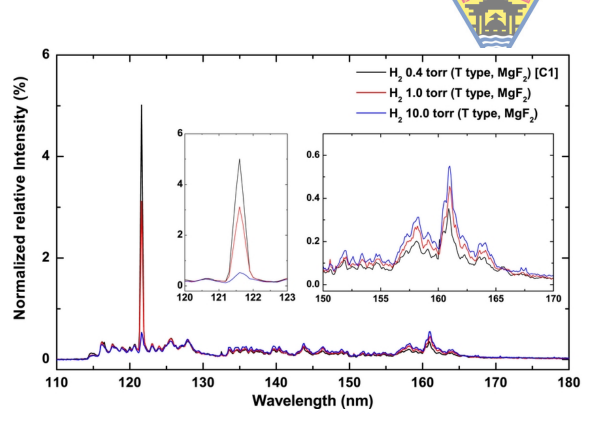


Figure 1.3: VUV spectra of MDHL (T-type geometry, 110-180 nm) with different H₂ pressure inside the lamp(Quoted from Chen et al. 2014).

 $\rm H_2$ with He or Ar etc), pressure of $\rm H_2$ and lamp geometry. Throughout those configurations stated there, we adopted 0.4 torr molecular hydrogen and T-type MDHL that produces VUV irradiation at 114-170 nm with 19.1 % of Ly- α and a mean photon energy of 9.27 eV. The photon flux is 6.4×10^{13} photons $cm^{-2}s^{-1}$ at sample position.

1.1.3 Extreme EUV source

To simulate the solar EUV irradiation reflected by IPM on both Charon and interstellar ices, we use the HF-CGM high – flux beam line of the National Synchrotron Radiation Research Center in Hsinchu, Taiwan. It provides a continuum EUV to VUV photons from 4 to 40 eV. The continuum is separated into monochromatic 30.4nm photons with a six-meter cylindrical grating monochrometer with an incident angle of 70 degrees. With the help of a movable entrance slit and movable curved exit slit, the energy resolving power can reach around 3×10^4 at 40 eV for grating 1600 l/mm with both slits movable and set opening to $10~\mu m$ [?]. Similar to VUV irradiation provided by MDHL, the light intensity was monitored by the same nickel mesh with photoelectric

efficiency obtained by SXUV 100 photodiode calibrated by NIST. With the known photoelectric efficiency, the flux of monochromatic 30.4nm is measured to be 2.15×10^{14} photons $s^{-1}cm^{-2}$ with a spot size of 1 cm

1.2 Experimental Protocol

In this section, we will briefly introduce the procedures of how we performed our experiments. It is divided into four parts, preparation and cooling, deposition, irradiation and warmup.

Preparation of experiments and cooling

Before any of experiment is done, we bake our system at 100 oC for 48 hours to reduce the contamination of water and residue gases as much as possible. It was cooled to room temperature that the background pressure can reach routinely at 1×10^{-10} torr. The gasline were connected with the regulators of the gas tanks and bake to 100 °C and pumped by molecularturbo pump for two days before any experiment were done. Also, The water sample has been freeze thaw several times by liquid nitrogen until there is no pressure increase recorded by baratron when water is freezed. Before cooling the substrate to cryogenic temperature, we took an IR spectrum and started the monitoring of residue gases by QMS in order to compare the residue molecules and to verify any possible contaminations in the main chamber. We then start the cooling process thanks to the closed-cycle He cryostat.

Deposition

The gas mixtures are pre-mixed in our gasline system introduced in section 1.1.1. We used a leak valve to condense the gas from the stainless steel bottles onto pre-cooled KBr substrate at 14 K, which monitored by Fourier transformed Infra-red spectroscopy (FTIR) and Quadrupole mass spectrometer (QMS) during deposition. The pressure of deposition is fixed to 1×10^{-8} torr that the deposition rate is $4 \times 10^{16} moleculescm^{-2} min^{-1}$. After deposition, we placed the ice mixture at 14 K for 60 minutes and to allow pumping of residue gas, until pressure of the main chamber reduce back to its base pressure to simulate the interstellar environment before irradiation.



Photon Irradiation

The total irradiation time is 270 to 450 minutes depend on experiment configurations; with time intervals varies from 2 to 30 minutes. After each irradiation, we waited for 10 minutes allowing pumping out of the photodesorpted gas molecules. During irradiation, the photon flux is monitored by a nickel mesh. After Irradiation, we place the sample for 30 minutes to observe if any thermal reaction was conducted.

Warmup

We use 1 K/min to warmup the substrate to 300 K to demonstrate effects of a new born star nearby an interstellar cloud. During warmup, we record the QMS from 1 to 100 a.m.u. to observe if there are low quantity of higher mass product formed during irradiation.

1.3 Infra-red spectroscopy and the Beer's Law

We used infra-red spectroscopy extensively in chapter 3 and 4, it is a powerful tool in studying molecular interactions during irradiation and warmup. We choose infra-red rather than Ramen spectroscopy because infra-red has lower energy that it would not change the structure of the ice mixture nor breaking any of the bonds. With different vibration modes, the energy absorbed by molecules are quantized. With the energy of absorption bands in infra-red spectrum, we may identify the functional group of the species. To simply classify, molecules can have, from less energetic, translational, rotational and vibrational motions. Generally, vibrational motions can be divided into stretching and bending. Stretching needs more energy than bending. For stretching, there exist Symmetric and Asymmetric stretching, while bending can be divided into In-plane Scissoring, rocking and out of plane Wagging and Twisting (Figure 1.4).

By Beer's Law, we may calculate the column density of the molecule with its functional groups, which are used to plot figures in chapter 3 and 4. Beer Lambert's Law suggest that when light passes through a



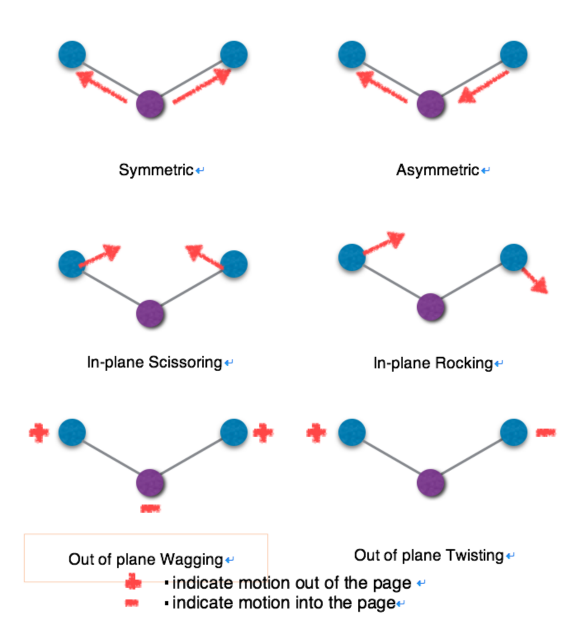


Figure 1.4: Different vibrational modes of a three atom molecule.

medium, amount of light absorbed is proportional to density and path length of the medium. Assume the known intensity beam $I_0(\nu)$ passes through the medium and beam intensity become $I(\nu)$. The transmittance $T(\nu)$ is defined by equation 1.1.

$$T(\nu) = \frac{I(\nu)}{I_0(\nu)} \tag{1.1}$$

Also, the absorbance $a(\nu)$ is defined by equation 1.2.

$$a(\nu) = -\ln T(\nu) = -\ln \frac{I(\nu)}{I_0(\nu)} = nl\sigma(\nu)$$
 (1.2)

where n is number density (molecules/cm³), l is the path length (cm), $\sigma(\nu)$ is the cross-section (cm²/molecule) of corresponding frequency ν . This equation is known as Lambert Beer's Law.

As the ice mixture in our thesis are at 14K, the peaks of absorbance are often a broadband due to coupling between neighbor molecules. Therefore, we can integrate the whole band of the peak equation 1.2 with respect to frequency and use the absorbance strength (A value) in literatures to calculate the column densities N of the ices by equation 1.3.

$$N = \frac{\int a(\nu) d\nu}{A(\nu)} \tag{1.3}$$

where N is the column density (molecule cm⁻²), $A(\nu)$ is the absorbance strength (cm molecule⁻¹).

1.4 Reaction Rate Laws

In this section, we will introduce rate reaction of a consecutive reaction and the concept of pseudo first order which we used to fit our reaction product against irradiation time. The rate of a chemical reaction is the relation between change in concentration of a substance per unit of time. i.e. For a balanced chemical reaction, $A \to 2B$, the rate of reaction is $-\frac{\Delta[A]}{\Delta t}$. The formation rate of B is 2 times destruction rate of A.

To determine the order of a reaction, we can only determine it experimentally. For a zero order reaction, the rate is not depending on any reactant that it is a constant. The rate $= -\frac{\Delta[R]}{\Delta t} = k[R]^0$. By calculus, $[R]_0 - [R]_t = kt$.

For a first order reaction, rate $=-\frac{\Delta[R]}{\Delta t}=k[R]$. By calculus, $\ln[R]_t=-kt+\ln[R]_0$.

For a second order reaction, rate $=-\frac{\Delta[R]}{\Delta t}=k[R]^2$. By calculus, $\frac{1}{[R]_t}-\frac{1}{[R]_0}=kt$.

Hence, if we get a straight line in a time versus concentration plot, it is a zeroth order reaction, similarly, in first order reactions, we get proportional relationships in time versus $\ln[R]$.

In a reaction with one reactant in excess, the rate of reaction is called pseudo first order reaction where pseudo means pretended. For $A+B \to C$, rate = k[A][B]. As $[B]_0 \gg [A]_0$, change of [B] is negligible that $[B] \sim [B]_0$. Therefore, [B] is assumed to be a constant and included in the rate constant k.

For a consecutive reaction equation, which we used to fit our data points, where $A \to B \to C$ that the produced product will not convert back as reactant. A simple example is radioactive decay. At t = 0, $[A] = [A]_0$, [B] = 0, [C] = 0 and at all times, $[A] + [B] + [C] = [A]_0$. The rate equations are as follows:

$$-\frac{\Delta[A]}{\Delta t} = k_1[A] \tag{1.4}$$

$$-\frac{\Delta[B]}{\Delta t} = k_1[A] - k_2[B]$$
 (1.5)

$$-\frac{\Delta[C]}{\Delta t} = k_2[B] \tag{1.6}$$

By equation 1.4, we get

$$[A] = [A]_0 e^{-k_1 t} (1.7)$$

By substituting equation 1.7 into equation 1.5, we get

$$-\frac{\Delta[B]}{\Delta t} + k_2[B] = k_1[A]_0 e^{-k_1 t}$$
(1.8)



After solving the differential equation 1.8 , we get

$$[B] = \frac{k_1}{k_2 - k_1} (e^{-k_1 t} - e^{-k_2 t}) [A]_0$$
 (1.9)

Finally, since $[C] = [A]_0 - [B] - [A]$, by equation 1.7 and 1.9, we get

$$[C] = \left(1 + \frac{k_1 e^{-k_2 t} - k_2 e^{-k_1 t}}{k_2 - k_1}\right) [A]_0 \tag{1.10}$$





2. Results and Discussions

According to the New Horizons team [?], CH_4 from Pluto may accumulate by cold-trapping, onto the surface of Charon. The amount of CH_4 varies along the surface of Charon because it depends on the length of time the temperature is below 25 K which in turns depends on diurnal motion and thermal inertia of Charon. With an axis tilted by 112 degrees from the ecliptic, higher concentration of CH_4 will be accumulated at the pole (see chapter ?? for details). In this chapter, we will investigate the following by infra-red spectroscopy: 1. The photo products produced by different concentration ratios of methane to ammonia, 2. the photo products produced by different photo sources (i.e. EUV and VUV) 3. the reaction mechanisms of each main products and 4. the functional group of tholin formed by irradiation of VUV, EUV on different configurations of CH_4+NH_3 ice mixtures (the result is compared with the residues on Titan produced by Imanaka et al. [?]

2.1 The infra-red spectrums and peaks identification

We scanned the IR spectrum before and after deposition and plotted the plot the corresponding absorbance of the ice mixtures. Figure 2.1 is a plot of the absorbance of the CH₄+NH₃ ice mixtures in different concentration ratios: 1:20, 1:10, 1:5 and 3:2 (arrangedfrom top to bottom). We labelled the peaks used in column density calculation by dotted lines in the graph. Main products we have detected are C₂H₆, CN⁻ and C₃H₈. The peak positions for substance identification are listed in Table 2.1. After identification of the products, we will look into each main products individually.

To calculate the column density, we integrate the area under graph



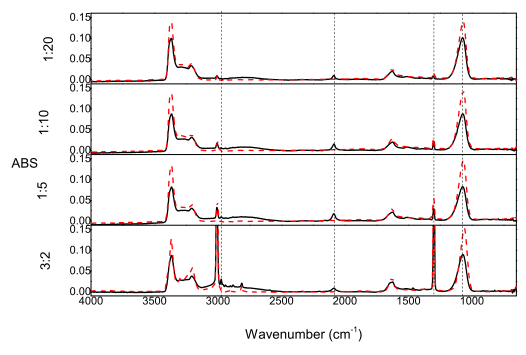


Figure 2.1: The the infra-red spectrum of $\mathrm{CH_4} + \mathrm{NH_3}$ ice mixtures before irradiation (dashed) and VUV irradiated ice mixtures provided by MDHL.

Table 2.1: The peak positions of identified substances after irradiation in different configurations of ice mixtures.

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Literti	ure assignments	CH	$I_4+\mathrm{NH_3}$ ra	atio (MDE	IL)	
Wavenumber	Carrier	1:5	1:10	1:20	3:2	Ref.
$({\rm cm}^{-1})$		$({\rm cm}^{-1})$	$({\rm cm}^{-1})$	$({\rm cm}^{-1})$	$({\rm cm}^{-1})$	
3375	$\nu_3 (\mathrm{NH_3})$	3366	3366	3369	3367	1
3290	$2\nu_4 (\mathrm{NH_3})$	-	-	-	-	1
3210	$\nu_1 \left(\mathrm{NH_3} \right)$	3207	3208	3210	3205	1
3011	$\nu_3 (\mathrm{CH_4})$	-	-	-	-	2
2972	$\nu_{10} \left(\mathrm{C_2H_6} \right)$	2975	-	-	2975	3
2960	$\mathrm{C_3H_8}$	-	-	-	2960	7
2941	$\nu_8 + \nu_1 1 (C_2 H_6)$	2940	-	-	2940	3
2904	$\nu_1 (\mathrm{CH_4})$	2901	-	-	2901	5
2879	$\nu_5 (\mathrm{C_2H_6})$	2882	2883	-	2882	3
2814	$\nu_2 + \nu_4 (\text{CH}_4)$	-	-	-	2815	5
2083	$\nu (\mathrm{CN^{-}})$	2088	2087	2088	2088	2
1625	$\nu_4 (\mathrm{NH_3})$	1625	1625	1626	1631	1
1514	$\delta (\mathrm{NH_2})$	1509	1507	1505	1511	6
1465 - 1440	deform CH ₂ scissor	1461	-	-	1463	3,4
1390-1370	CH_3 sym deform	1394	1394	1394	1372	4
1298	$\nu_4 (\mathrm{CH_4})$	1301	1302	1305	1299	2
1075	$\nu_2 (\mathrm{NH_3})$	1073	1072	1072	1072	1
820	$\nu_1 2 \; ({\rm C_2 H_6})$	-	-	-	820	3
D -f 1	D4 -1 9000 [2]	0 1/	1 TT 1	2002	[9] o TZ:	4 .1

Reference: 1. Bossa et al. 2008 [?] 2. Moore and Hudson 2003 [?] 3. Kim et al. 2010 [?] 4. Socrates 2001 [?] 5. Bennet and Kaiser 2007 [?] 6. Zheng et al. 2008 [?] 7. Hudson and Moore 2004 [?]

and divided by the absorption strength presented in table 3.2. We aware that there is an average error in absorption strengths of no more than 10 % when the pure ice is diluted in N_2 and H_2O [?]. In our case, absorption strengths changes after CH₄ and NH₃ are mixed. For example, according to d' Hendecourt and Allamandola [?], the band of NH_3 located at 1070 cm⁻¹ would not change much (from 1.1×10^{-17} to 1.2×10^{-17}) when excess water is added to pure NH₃. For the case of $\mathrm{CN^-}$, we know that $\mathrm{CN^-}$ has a bond order =3 from its molecular orbitals. CN⁻ which is different from CN (bond order 2.5). CN stretching is very sensitive to the matrix environment. It can change by factor of 2 in amino acetonitrile and H_2O (1:3) [?]. However, CN is not inspected in this chapter. Therefore, we are justified to use the same absorption strength throughout our discussion to estimate the column density of each species and how the absorption area changes with concentration ratios of ice mixtures and photon energy. Here, we adopt the absorption strengths stated in Table 2.2

Table 2.2: The strength of absorbance adopted in this thesis measured in literatures of pure ice samples

Wavenumber (cm ⁻¹)	Assignment	Vibration	FWHM	A value ($\times 10^{-17}$)	Reference
2976	C_2H_6	$-\mathrm{CH}_3$	-	1.05	2
2960	C_3H_8	$-CH_2$ -	-	2.58	2
2086	CN^-	CN	-	1.8	3
1297	CH_4	CH deformation	8	0.61	1
1070	NH_3	"umbrella mode"	68	1.7	1

Reference: 1. d'Hendecourt and Allamandola (1986)[?] 2. Moore and Hudson (1998)[?] 3. Noble et al. (2013) [?]

2.2 Reaction mechanisms

2.2.1 C_2H_6

The assignment of C_2H_6 is confirmed by several bands listed in table 2.1. Figure 2.2 is a zoomed view of figure 2.1. The absorption peak located at 2075 cm⁻¹ corresponds to the strongest vibration of C_2H_6 . The formation of C_2H_6 in astrophysical environment is mainly through a combination with 2 CH₃ radicals [?]:

$$CH_4 + hv \rightarrow CH_3$$
 (2.1)



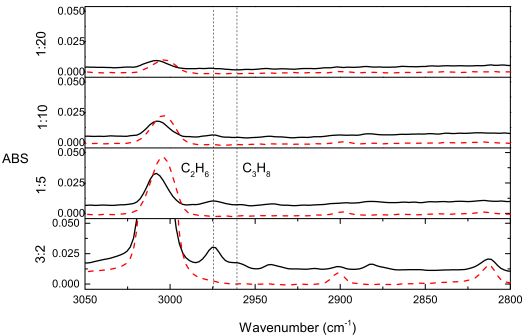


Figure 2.2: The the infra-red spectrum of $CH_4 + NH_3$ ice mixtures of C_2H_6 and C_3H_8 before irradiation (dashed) and VUV irradiated ice mixtures provided by MDHL.

$$2CH_3 \to C_2H_6 \tag{2.2}$$

The energy required to produce 1 $\rm CH_3$ radical from $\rm CH_4$ is 4.42 eV. Recombination of 2 $\rm CH_3$ radicals to form $\rm C_2H_6$ releases 3.74 eV. The process in 2.2 is a no-barrier exothermic process. Note that $\rm C_2H_6$ is not detected in $\rm CH_4$ to $\rm NH_3{=}1{:}20$ ice mixtures. Figure 2.3 shows the temporal formation column density of $\rm C_2H_6$ in different configurations of irradiated ice mixtures. As the formation only depends on $\rm CH_4$, we may use first order kinetics equation to fit the column density versus photon dose.

$$[A] = [A]_0(1 - e^{-k_1 t}) (2.3)$$

The fitting results are shown in table 2.3.

Table 2.3: The fitting results of C_2H_6 by $[C_2H_6]=[C_2H_6](1-e^{-k_1t})$

Ratio of CH ₄ +NH ₃	A $(x10^{15} \text{ molecules cm}^{-2})$	$k (x10^{-17} \text{ photon}^{-1})$
1:10	2.90 ± 1.25	0.92 ± 0.15
1:5	4.16 ± 0.28	2.28 ± 0.28
3:2	19.2 ± 0.15	5.28 ± 0.25

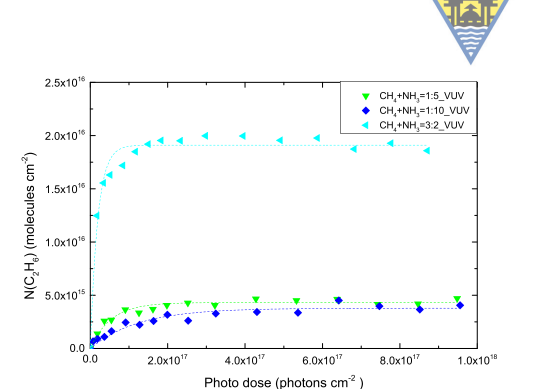


Figure 2.3: The column density of C2H6 during CH4 + NH3 ice mixtures irradiated by MDHL.

From table 2.3, the production rate is nearly proportional to the initial CH_4 concentration.

$2.2.2 C_3H_8$

The peak positioned at 2960 cm⁻¹ belongs to $-CH_2$ - so we assigned that as C_3H_8 , as the shortest carbon chain. The signal to noise ratio in $CH_4+NH_3=1:10$ is poor that we can not quantize the amount of C_3H_8 (figure 2.2).

It is a secondary product formed by a combination of either $C_2H_6 + CH_2$ (equation 2.4) or $C_2H_4 + CH_4$ (equation 2.5).

$$C_2H_6 + CH_2 \to C_3H_8$$
 (2.4)

$$C_2H_4 + CH_4 \to C_3H_8$$
 (2.5)

By modern peak fitting method, we deconvoluted the overlapped C_2H_6 and C_3H_8 into two gaussians.

2.2.3 CN^{-}

From infra-red absorption spectrum (figure 2.4) and their positions, we assigned the peak 2086 cm⁻¹ to CN⁻ but not a combination of HCN

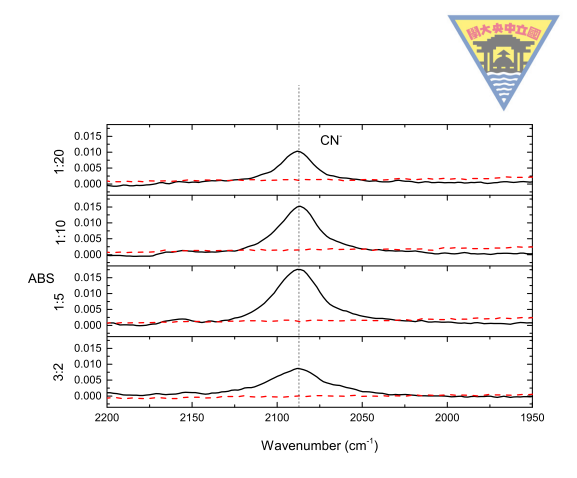


Figure 2.4: The the infra-red spectrum of $CH_4 + NH_3$ ice mixtures of C_2H_6 and C_3H_8 before irradiation (dashed) and VUV irradiated ice mixtures provided by MDHL.

and $\rm CN^-$. The assignment is based on a absence in CN bending mode at 848 cm⁻¹. In the case $\rm CH_4 + NH_3 = 3:2$, we may observe a peak located at 820 cm⁻¹, which is with a FWHM half of HCN and it is eliminated at 50 K during the warm-up phase. Since 50 K is the desorbing temperature of $\rm C_2H_6$ and the peak position is the close to v12 mode of $\rm C_2H_6$, we believe that the 820 cm⁻¹ peak is contributed by $\rm C_2H_6$. Therefore, we may assign our peak located at 2086 cm⁻¹ as purely CN⁻.

The formation mechanism of CN⁻ at low temperature was first suggested by Kim and Kaiser [?] to be two step reaction mechanism with methylamine as intermediate. CH₄ and NH₃ irradiated by photon to become CH₃ and NH₂ radical (figure 2.5, followed by propagation and recombination of radicals becoming CH₃NH₂ and dehydrogenation and acid-base reaction to form CN⁻. Although Kim and Kaiser [?] used 1.5 keV electron as energy source to simulate the cosmic ray induced photochemistry, this formation mechanism also applies in our photon



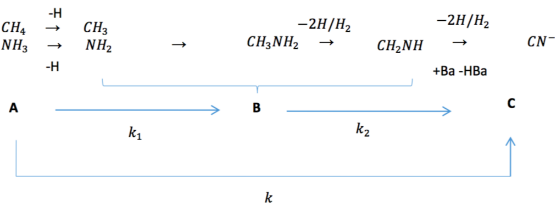


Figure 2.5: The formation mechanism of CN⁻ proposed by Kim and Kaiser(2011).

irradiation experiments because we can also detect the methylamine during our warm-up phase. The ion fragment with m/z=31 is assigned as $CH_3NH_2^+$ and detectable in all ratios of our CH_4+NH_3 experiments (figure 2.6).

By the deviation performed in section 1.4, we have a rate equation for consecutive reactions 1.10. With one of the reactant larger than another, we applied the pseudo first order assumption. With equation 1.10, we fitted the formation of CN^- (figure 2.7) and found that one of the rate constant is always larger than the other in all of the ratios. The fitting results are averaged by more than two experiments and are shown in table 2.6. The results of Kim and Kaiser is also listed into the table, they could observe a two-step reaction mechanism in production of CN^- in CH_4+NH_3 (3:1) experiments with electron current 0.1 μ A. However, when they increased the electron flux to 1 μ A for irradiation $C_nH_{2n+2(n=1-6)}$ and NH_3 ice mixtures, they also observed a one-step reaction mechanism.

Table 2.4: The fitting results of CN⁻ by equation 1.10

Ratio of CH ₄ +NH ₃	A $(x10^{16} \text{ molecules cm}^{-2})$	$k_1 (x10^{-18} \text{ photon}^{-1})$	$k_2 \text{ (photon}^{-1})$	
1:20	4.75 ± 0.40	0.70 ± 0.09	>1	
1:10	4.51 ± 0.18	1.33 ± 0.13	>1	A
1:5	4.61 ± 0.18	1.93 ± 0.19	>1	
3:2	2.24 ± 0.03	8.21 ± 0.70	>1	

represents the amount of CN⁻ we may obtain when irradiated the ice for infinitely long.



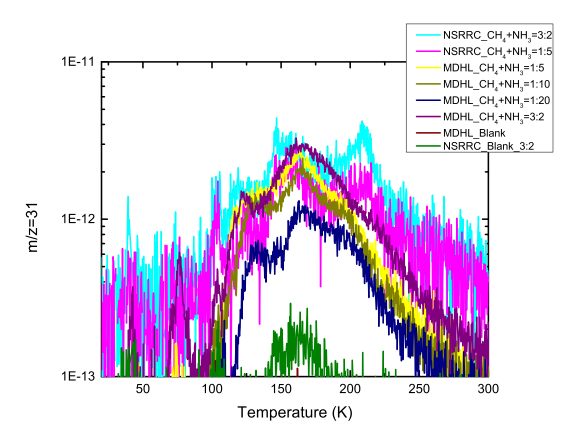


Figure 2.6: The m/z=31 detected by QMS during warm-up with heating rate 1 K/min in different configurations of ice mixtures.

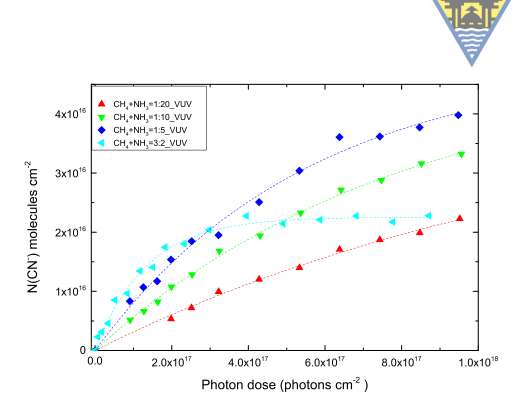


Figure 2.7: The column density of CN^- accumulated when different configurations of $CH_4 + NH_3$ ice mixtures are irradiated by VUV photons provided by MDHL. The dotted lines are fits of column densities by equation 1.10.

2.3 The Concentration Effect in formation of Cyanide ions and Ethane

2.3.1 Cyanide ion

From table 2.6, we may observe that the rate k_1 is nearly proportional to the concentration of CH_4 . As CH_4 to NH_3 ratio increases, more CH_4 are involved in CH_3 radical formation, thus there are more CH_3 radicals to produce CH_3NH_2 intermediates.

In CH_4 to $NH_3 = 3:2$ ice mixtures, A is about half of that of the other ratios. The reduction is mainly because NH_2 (forming CH_3NH_2) has a competing relationship with CH_2 , CH_3 and C_2H_4 radicals (forming C_2H_6 and C_3H_8). This competition supresses the production of intermediate CH_3NH_2 , thus the formation of CN^- . Therefore, the yield of CN^- is the least in CH_4 to NH_3 ice mixture with ratio 3:2 while the yield of C_2H_6 is the greatest in the mixture with the same ratio(table 2.6), (table 2.3)

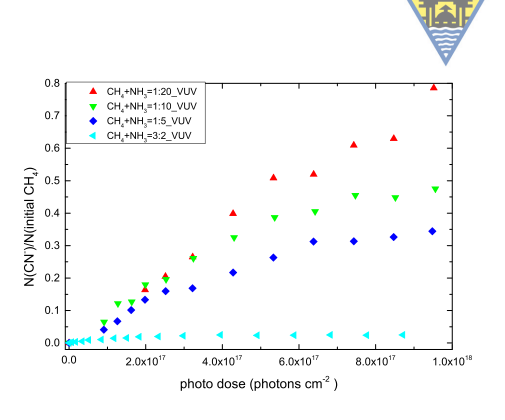


Figure 2.8: The column density of $\rm CN^-$ divided by initial $\rm CH_4$ accumulated when different configurations of $\rm CH_4 + NH_3$ ice mixtures are irradiated by VUV photons provided by MDHL.

Considering the normalized CN⁻ with respect to the initial CH₄(figure 2.8), the formation of CN⁻ is more efficient in low CH₄ concentration ice mixtures. At low CH₄ concentration, there are excess NH₃ which can aggregate mobile CH₃ radicals, preventing meeting another CH₃ radical or C₂H₄. Therefore the production of C₂H₆ is greatly suppressed and more CN⁻ will be produced.

2.3.2 Ethane

Considering the case of ratio of CN⁻ divided by C₂H₆, the formation of CN⁻ in ice mixtures with diluted CH₄ has more CN⁻ formed then C₂H₆. It is because ice mixtures with with higher concentrations in CH₄ is more effective for one CH₃ radical to combine with another CH₃ radical. On the contrast, CH₃ radicals formed in the ice mixtures with diluted CH₄ concentrations are aggregated by NH₃. Therefore, CN⁻ is less efficient to form in ice mixtures with excess NH₃.



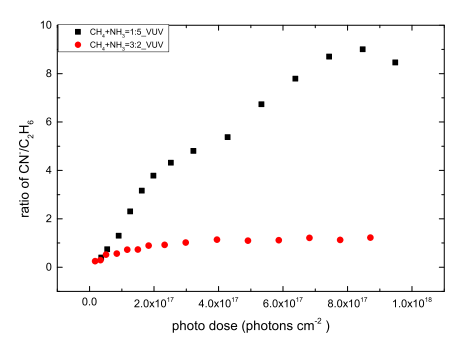


Figure 2.9: The column density of $\rm CN^-$ divided by $\rm C_2H_6$ accumulated when different configurations of $\rm CH_4 + NH_3$ ice mixtures are irradiated by VUV photons provided by MDHL.

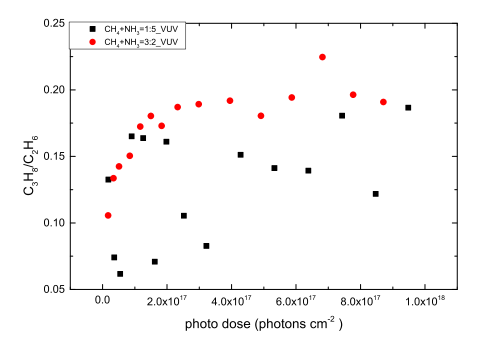


Figure 2.10: The column density of C_3H_8 divided by C_2H_6 accumulated when different configurations of $CH_4 + NH_3$ ice mixtures are irradiated by VUV photons provided by MDHL.



2.3.3 Propane

 C_3H_8 forms based on to the C_2H_6 2.10 is the plot with column densities of C_2H_6 divided by C_3H_8 . We may see that the ratio in CH_4+NH_3 =1:5 experiment is around 6 where CH_4+NH_3 =3:2 is around 3. This shows that the amount of C_3H_8 in CH_4+NH_3 =3:2 experiment is higher. It is rather difficult for C_3H_8 to form in CH_4+NH_3 = 1:5 experiments because NH_3 aggregated them. The formation of C_3H_8 in CH_4+NH_3 =1:5 and 3:2 experiments has given a reasonable explanation about why C_2H_6 formation is most efficient in CH_4+NH_3 =1:10 experiments.

2.4 Photon Energy Effect - EUV and VUV

According to Blanksby and Ellison [?], the dissociation energy for CH₄, becoming CH₃, CH₂ CH and C are 4.55, 4.79, 4.39 and 3.51 eV respectively at 298 K. Whereas dissociation energy for NH₃, becoming NH₂ is 4.67 eV at 298 K.

Considering our MDHL with average energy of 9.27 eV, all of the above fragments may exist either in the form of radicals or combined with other radicals to form heavier molecules in our ice mixtures. Although Increasing the photon energy does not create new fragmentation pathway, the choice of fragmentation pathways depends on photon energy.

Several gaseous state measurements also support this statement. First, Gans et al. (2011) [?] changed VUV photon wavelengths from 121.6 nm to 118.2 nm to dissociate the CH₄ molecules and ionize the fragments with the corresponding photon energy. Changing the output of the pulsed laser from 121.6 to 118.1 nm significantly changed the ratio of CH₃⁺ and CH₂⁺, produced from fragmentation, from 1: 1 to 1:2. This slight change of photon energy, from 10.2 eV to 10.4 eV has a significant change in the ratio between different pathways.

Second, an EUV fragmentation experiment done by Tsai et al. [?] used 30.4 nm to photo-dissociate CH_4 and tested it by timeĂŞ-ofĂŞ-flight mass spectrometer yields CH_3^+ : CH_2^+ : CH^+ : $C^+ = 1 : 0.32 : 0.118 : 0.0237$ (Tsai 1980). Consider the ratios of CH_3 to CH_2 radicals, it is around 3 to 1, which is in contrast to the experiment results of Gans

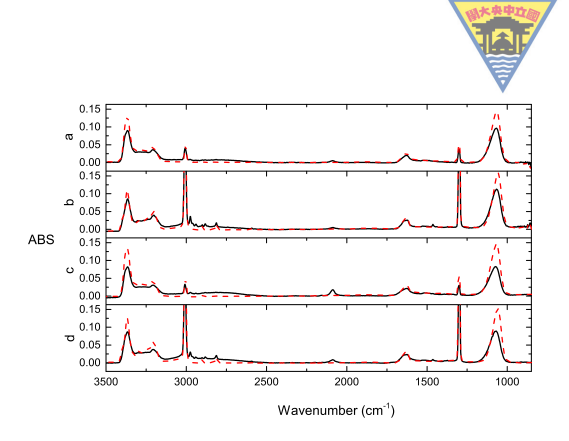


Figure 2.11: The the infra-red spectrum of $CH_4 + NH_3$ ice mixtures before irradiation (dashed) and VUV and EUV (solid) irradiated ice mixtures provided by MDHL. (a) and (b) are EUV irradiated $CH_4 + NH_3 = 1:5$ and 3:2 ice mixtures respectively, and (c) and (d) are VUV irradiated $CH_4 + NH_3 = 1:5$ and 3:2 ice mixtures respectively.

et al. (2011)[?]. Although both of them are gaseous state experimental results, it is uncertain that if increasing photon energy can produce more CH₂ radicals.

Thirdly, a group varies ratios of $CH_4 + NH_3$ mixtures and irradiate with far UV irradiation at 134 nm [?]. However, this group only used gas chromatography to analyse the final products and their reaction is carried in gas phase in room temperature. We aware that the VUV absorption spectra of CH_4 in solid phases is different from gaseous phases [?], so the exact photo dissociation fragmentation ratios by EUV nor VUV irradiations in astronomical environments are still unknown. It is worthwhile for us to perform the experiment by EUV irradiation to see if EUV irradiation can generate any new products on the surface of Charon, or any difference in yield. Despite the photon energy of our MDHL is enough to dissociate both the CH_4 and NH_3 molecules, we further increase photon energy to He II 30.4 nm to examine the differences in photo-products.

Table 2.5 shows the identified peaks of CH_4+NH_3 ice mixtures irradiated by VUV and EUV (30.4 nm) irradiated in IR spectra (figure 2.11).

Table 2.5: The peak positions of identified substances after VUV and EUV irradiations in different configurations of ice mixtures.

Literture assignments		CH ₄ +NH ₃ ratio (MDHL)		CH ₄ +NH ₃ ratio (30.4 nm)		
Wavenumber	Carrier	1:5	3:2	1:5	3:2	Ref.
(cm^{-1})		$({\rm cm}^{-1})$	$({\rm cm}^{-1})$	$({\rm cm}^{-1})$	(cm^{-1})	
3375	$\nu_3 (\mathrm{NH_3})$	3366	3367	3368	3368	1
3290	$2\nu_4 \text{ (NH}_3)$	-	-	-	-	1
3210	$\nu_1 (\mathrm{NH_3})$	3207	3205	3209	3205	1
3011	ν_3 (CH ₄)	-	-	-	-	2
2972	$\nu_{10} \; (\mathrm{C_2H_6})$	2975	$2975\ 2977$	2976		3
2960	C_3H_8	-	2960	-	2960	7
2941	$\nu_8 + \nu_1 1 \ (C_2 H_6)$	2940	2940	-	2942	3
2904	$\nu_1 (\mathrm{CH_4})$	2901	2901	2901	2901	5
2879	$\nu_5 (\mathrm{C_2H_6})$	2882	2882	-	2884	3
2814	$\nu_2 + \nu_4 (\text{CH}_4)$	-	2815	-	2813	5
2083	$\nu (\mathrm{CN}^-)$	2088	2088	2090	2089	2
1625	$\nu_4 (\mathrm{NH_3})$	1625	1631	1627	1631	1
1514	$\delta (\mathrm{NH_2})$	1509	1511	1509	1511	6
1465-1440	deform CH ₂ scissor	1461	1463	-	1465	3,4
1390-1370	CH_3 sym deform	1394	1372	-	1372	4
1298	$\nu_4 \ (\mathrm{CH_4})$	1301	1299	1303	1301	2
1075	$\nu_2 (\mathrm{NH_3})$	1073	1072	1070	1068	1
820	$\nu_1 2 (C_2 H_6)$	-	820	-	-	3

Reference: 1. Bossa et al. 2008 [?] 2. Moore and Hudson 2003 [?] 3. Kim et al. 2010 [?] 4. Socrates 2001 [?] 5. Bennet and Kaiser 2007 [?] 6. Zheng et al. 2008 [?] 7. Hudson and Moore 2004 [?]

Considering the formation mechanisms of C_2H_6 and C_3H_8 , equation (2.2 and 2.4), when MDHL VUV irradiation is replaced by He II 30.4 nm monochromatic light, the ratio of C_2H_6 to C_3H_8 in CH_4 to $NH_3=3:2$ ice mixtures irradiated by VUV irradiation is lower under EUV irradiation than that under EUV provided by NSRRC (figure 2.12). There are two possible explanations. First, different photon energies flavour different CH_4 fragmentation pathway and less C_3H_8 is produced with EUV photons. Second, the efficiency of CH_4 fragmentation is greatly reduced under EUV irradiation and the density of CH_3 radicals are much lower than that in the case of VUV irradiation provided by the MDHL. We lengthen the time of EUV irradiation on our ice mixtures until the total number of destructed CH_4 is similar to that in VUV irradiation experiments done with MDHL. The averages of ratios of $C_2H_6:C_3H_8$ of the last 7 irradiations before terminating irradiations are 3.53 in VUV and



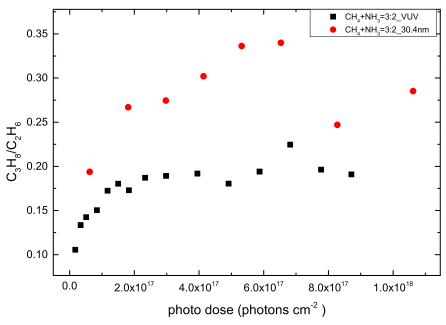


Figure 2.12: The column density of C_3H_8 divided by C_2H_6 accumulated when different configurations of $CH_4 + NH_3$ ice mixtures are irradiated by VUV and EUV photons

3.66 in EUV. The result supports the latter explanation. From figure 2.13, The reduction of CH_4 is $6.06\pm$ times slower in EUV experiments than VUV experiments while the reduction of NH_3 is 3.19 ± 0.12 times slower. Therefore, the destruction cross-section of CH_4 and NH_3 ice has a 6.06 ± 0.07 and 3.19 ± 0.12 times lower in 30.4 nm than in 121.6 nm.

Figure 2.12 shows the column densities of C_2H_6 divided by C_3H_8 after $CH_4 + NH_3 = 3:2$ ice mixtures are irradiated by VUV irradiation and He II monochromatic light.

From 2.12, we may observe that more C_3H_8 is produced by 30.4nm photons than by VUV photons. Recall the formation mechanism of C_3H_8 (equation 2.5), CH_2 and C_2H_4 radicals are escential in producing C_3H_8 . This increase production in C_3H_8 may be caused by the increase in CH_2 radicals during fragmentation of CH_4 . This result is similar to the findings of Gans et al. (2011)[?], the ratio of CH_2 radicals increases from 0.3 to 0.48 when photon energy increases from 121.6 nm to 118.2 nm in their pulsed laser experiments.

Apart from C₂H₆ and C₃H₈, are there any difference in CN⁻ pro-



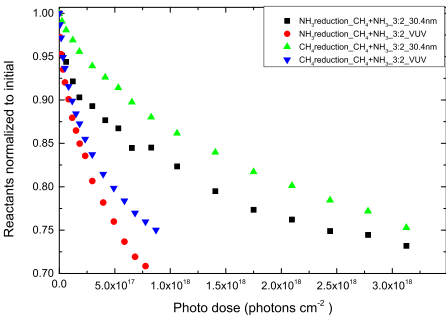


Figure 2.13: The normalized reduction of CH_4 and NH_3 in $CH_4 + NH_3$ ice mixtures irradiated by VUV and EUV photons

duction? Figure 2.14 shows the accumulated column densities of CN⁻ generated by irradiation of CH₄+NH₃ ice mixtures by MDHL and 30.4 nm monochromatic light. The fitting results are shown in Table 3.6. The rate constants forming CN⁻ is 3.06 to 4.13 times larger in CH₄+NH₃ = 1:5 and 3:2 irradiated by MDHL than irradiated by 30.4 nm monochromatic light respectively. From figure 2.13, the CH₄ reduction in NSRRC is 6.06±0.07 times slower. With the rate constants of CN⁻ only 3.06 to 4.13 times smaller, the 6 times slower in CH₄ reduction and 3 times slower in CN⁻ formation give rise to a similarity of reduced NH₃ destruction cross-section and reduced rate in CN⁻ production in EUV irradiation experiments. Therefore, we may conclude that the reduction in CN⁻ formation rate by 30.4nm EUV irradiation is mainly due to the decreased NH₃ destruction cross-sections.

2.5 Residues

The residues we studied are the accumulated residues onto the substrate. We do not understand are there any interaction between residues and the ice mixtures. However, we may know what is the change of



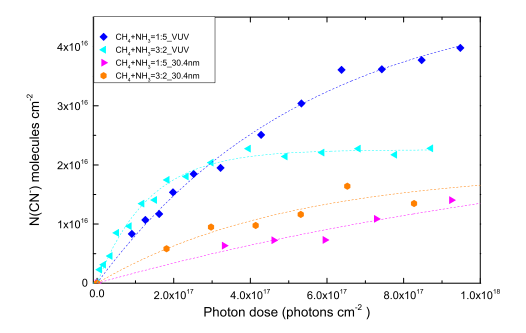


Figure 2.14: The column densities of $\rm CN^-$ generated by irradiation of $\rm CH_4+NH_3$ ice mixtures by MDHL and 30.4 nm monochromatic light.

Table 2.6: The fitting results of CN^- by equation 1.10

Light source	Ratio of CH ₄ +NH ₃	A $(x10^{16} \text{ molecules cm}^{-2})$	$k_1 (x10^{-18} \text{ photon}^{-1})$	$k_2 \text{ (photon}^{-1})$
VUV	1:5	4.61 ± 0.18	1.93 ± 0.19	>1
MDHL	3:2	2.24 ± 0.03	8.21 ± 0.70	>1
EUV	1:5	2.89 ± 1.29	0.63 ± 0.37	>1
$30.4\mathrm{nm}$	3:2	2.24 ± 0.03	1.92 ± 1.99	>1

Fitting result of figure 2.14 with pseudo first order equation $[CN^-]=A(1-e^{-kx})$. These fitting results of MDHL experiments are an average of at least 2 experiments with the same circumstances. In the expression, A represents the column density when x, the photon dose, becomes infinitely large and k is the rate constant.

residues when we change the ratio of the CH_4+NH_3 from CH_4 dominating to NH_3 dominating. Figure 2.15 is a comparison of $CH_4+NH_3=3:2$ after VUV experiments, residues accumulated after EUV exposure of $CH_4+NH_3=3:2$ ice mixtures and the plasma experiment done by Imanaka et al. (2004)[?]. The residues in ammonia dominated ice mixtures cannot be detected after consecutive experiments. There are no differences between EUV accumulated residues and VUV accumulated residues in $CH_4+NH_3=3:2$ ice mixtues. The main differences between plasma experiments of N_2+CH_4 (9:1) done at 2300 Pa. by Imanaka et al. (2004)[?] and our experiments is the peaks located around 2090 cm⁻¹.

Why we use different initial reactants, replacing N_2 by NH_3 but we may get similar residues? The similarities during formation of atomic nitrogens when breaking N_2 bonds in nitrogen and NH bonds in ammonia give rise to this result. When photon energy is enough to break both NH bond and N_2 bond, similar experimental residues forms. Our results implies that the residues formed on Charon is similar to what we found on Titan, although their formation environments differs from gaseous phase with N_2 dominating to solid phase with NH_3 .

After $CH_4+NH_3=1:5$, 1:10 and 1:20 experiments, we notice that two new bonds are formed. One at 1721 cm^{-1} and another at 1286 cm^{-1} . These two peaks are due to MCT detector self-contamination. When we stopped adding liquid nitrogen, molecules stick onto MCT detector will be free out. They sticked onto our detector and hence produced these two peaks. Hence, we may conclude that the residues produced by CH_4+NH_3 with different ratios are the same.

2.6 Conclusion

The main product of VUV and EUV irradiated CH_4+NH_3 ice mixtures are C_2H_6 and CN^- . C_3H_8 is also produced by C_2H_6 or C_2H_2 . We did several investigations towards CH_4+NH_3 ice mixtures. First, by changing ratio of CH_4 and NH_3 in the ice mixture, CN^- production is more effective in NH_3 dominated ice mixtures where C_2H_6 dominates in CH_4 dominated ice mixtures. Second, by changing the photon source to EUV irradiation, the yield of C_3H_8 increases. The effective formation of



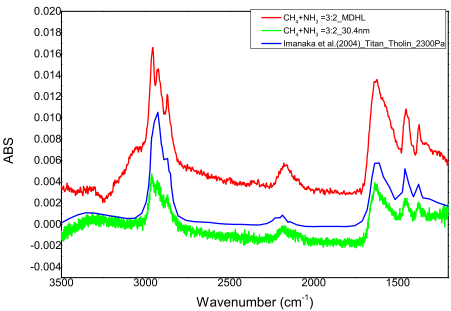


Figure 2.15: The IR spectrum of residues in after $CH_4+NH_3=3:2$ experiments and the accumulate residues after MDHL experiments and NSRRC experiments.

C₃H₈ was not produced by C₂H₆ but by C₂H₂ and CH₄ because the ratio of C₃H₈: C₂H₆ increases. This suggests that the CH₂ or CH fragmentation from CH₄ increases when photon energy increases. By studying the production efficiencies, the difference in photo-production yield is mainly caused by the reduction in photo-destruction cross-section in the reactants. Thirdly, by comparison with electron irradiation experiments, electron irradiation has a smaller absorption cross-sections, the percentage of yield is also smaller than VUV irradiated ice mixtures with similar ice thicknesses. Finally, we compared our residues obtained with laboratory produced Taitan tholins, the similar infra-red spectrum shows a similar functional groups in residues. Our result implies that the tholin on Charon should be similar to the tholin formed on Titan.





3. Astrophysical Implications

In this text, we look at effects The main source to irradiating the dark side of Charon is Ly α reflected by interplanetary medium [?]. Other sources included energetic ions in solar wind, which mainly consist of H⁺, He⁺⁺, He⁺⁺ and O²⁺ etc. originated from solar corona or IPM. These ions also reflect solar irradiation to the dark side of Charon. Among sources focused on He II irradiation as it is 3 toĂŞ 20 times more intense than He I during a solar flare. As the intensity varies with solar activities, it is difficult to estimate the dose onto Charon. Besides, electronic flux is also present in solar wind but it is one order of magnitude lower than proton flux. The flux for energetic electrons observed at the 1 A. U. position is available (http://www.swpc.noaa.gov/products/goeselectron-flux). Although electron flux is much less important than Ly- α , and their flux varies, we also compare the electron irradiation experiment done by Karie electrons.

3.1 Cyanide ion produced by photon source and electron source

We study the ice mixtures in which CH_4 is dominated and compare the efficiencies in CN^- formation by electrons and VUV irradiations. Comparing our CN^- obtained after infinitely long exposure, 13-16 ML of CN^- was obtained by electron irradiation depending on which equation they choose to fit. In our MDHL experiments, we have 14.8 ML of CN^- . However, Kim and Kaiser (2011) adopted the CN^- absorption coefficient measured by Georgieva and Velcheva (2006) to be 3.7×10^{-18} cm molecule⁻¹, which is 4.86 times smaller. We do not adopt this absorption coefficient because it violates the carbon balance that number of CN^- produced will be larger than CH_4 consumption. If we adopted

the same absorption coefficient, the production yield of CN should be multiplied by 4.86. Therefore, our yield is 72 ML of CN $^-$. Regarding percentage of NH $_3$ (limiting reactant), Kim and Kaiser has 5 - 6 % yield where we have 12 % yield if we adopted the same absorption coefficients.

Note that Kim and Kaiser kim have ice thickness (CH₄ = 610 ML, NH₃= 260 ML) while we have thicker ices (CH₄ = 900 ML, NH₃= 600 ML). Therefore, we calculate the percentage of photons absorbed by CH₄+NH₃ ice mixtures under VUV irradiation. Substituting cross-sections measured by Cruz-Diaz et al. (2014) and the spectrum of our MDHL and substitute them into Beer's law, CH₄+NH₃ = 3:2 ice mixtures can absorb more than 99 % of light when thickness of CH4 and NH3 equals

In electron irradiation experiments of Kim and Kaiser (2011)[?], the energy transferred to $CH_4 + NH_3$ ice mixtures is by linear electron transfer (LET) that 1.3 eV molecule⁻¹ was absorbed by the ice in 90 minutes. They get flattened at 20 minutes' irradiation, with fluence of 2.0×10^{14} electrons cm⁻². While we got flattened at a dose of 3×10^{17} photons cm⁻². Considering the energy of their electron (1.5 keV) and energy of our photons, they got flattened at 3×10^{17} eV cm⁻² while we get flattened at 27.81×10^{17} eV cm⁻². Comparing these energy doses, less electrons are needed to flatten the formation of CN^- .

To conclude, electron irradiation has a smaller absorption crosssections, the percentage of yield is also smaller than VUV irradiated ice mixtures with similar ice thicknesses.