THE STUDY OF INTERSTELLAR CHEMISTRY THROUGH MID-INFRARED SPECTROSCOPY

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ABSTRACT

We present mid-infrared spectra of various hydrocarbons toward the hot core associated with the massive protostar NGC 7538 IRS 1. Because high abundances of fully hydrogenated molecules and saturated molecules are characteristic of hot cores, they provide laboratories in which to study interstellar chemistry. Through the study of C$_2$H$_2$, HCN, NH$_3$, CH$_4$, and CH$_3$, we can constrain chemical models. We derive abundances for C$_2$H$_2$ and HCN and compare to model predictions. The observations were taken with the Texas Echelon Cross Echelle Spectrograph (TEXES).

Key Words: ISM: MOLECULES — ISM: ABUNDANCES — ISM: CLOUDS — ISM: MOLECULES

1. INTRODUCTION

The chemistry in star forming regions can be a useful tool in determining a sequence of massive star formation (e.g. van Dishoeck & van der Tak 2000). While there is a formation sequence for low-mass stars (Adams et al. 1987), such a sequence has not yet been determined for high-mass stars because massive stars can reach the main sequence while still embedded in dense molecular gas. Icy grain mantles are formed when molecules freeze out during the prestellar and collapse phases. Ices then evaporate when the protostar heats its surroundings. These newly evaporated molecules drive a rich chemistry in compact, optically thick hot cores. Hot cores have temperatures $\geq 100$ K, densities $\geq 10^7$ cm$^{-3}$, and diameters $\leq 0.1$ pc (Kurtz et al. 2000). Hot cores contain high abundances of fully hydrogenated molecules such as H$_2$O, NH$_3$, H$_2$S, as well as complex organics such as CH$_3$OH, CH$_3$CN, and CH$_3$OCH$_3$.

Because of the large extinction toward regions of massive star formation, most studies are done with millimeter or radio (hereafter simply radio) spectroscopy. Along with radio spectroscopy, mid-infrared absorption spectroscopy is a powerful probe of physical conditions and chemical abundances in molecular clouds. In infrared absorption spectroscopy, the spatial resolution is set by the size of the background source, whereas, in radio spectroscopy, the spatial resolution is set by the size of the beam. In our study, the massive protostar provides a background source and we can study the material close to the star. Moreover, mid-infrared spectroscopy allows for the study of molecules without a dipole moment such as acetylene, C$_2$H$_2$, and methane, CH$_4$ (Lacy et al. 1989b, Evans et al. 1991, Carr et al. 1995), which do not have rotational transitions. Previous mid-infrared studies have had low to moderate spectral resolution. For example,
the Short Wavelength Spectrometer (SWS) on board the Infrared Space Observatory (ISO) had resolving power, R, ~ 2000 and from Lacy et al. had R ~ 10^4. Now, a new instrument, the Texas Echelon Cross Echelle Spectrograph (TEXES) (Lacy et al. 2002), is available with resolving power R ≈ 80000, or δν ~ 4 km s^{-1} at 13μm. TEXES allows for a new class of observations and addresses long-standing questions such as velocity structure. NGC 7538 IRS 1, the brightest mid-infrared source (L = 1.3 × 10^5 L_⊙) in the NGC 7538 region, is a massive protostar with an ultra-compact HII region and possibly a hot core. Using ISO-SWS, Lahuis & van Dishoeck (2000) observed the Q-branches of HCN and C_2H_2 toward NGC 7538 IRS 1. The derived excitation temperatures are 500 K for C_2H_2 and 600 K for HCN. In this paper, we derive abundances for C_2H_2 and HCN and compare to chemical model predictions.

2. OBSERVATIONS

We have observed rovibrational lines for the following molecules toward NGC 7538 IRS 1: HCN, C_2H_2, CH_4, NH_3, and CH_3. Observations were made with TEXES (Lacy et al. 2002) at the NASA Infrared Telescope Facility in 2001 June, 2001 November, and 2002 September. The typical line width is ~ 5 km s^{-1} for both C_2H_2 and HCN lines. The paper with the complete data set is in preparation. Only the C_2H_2 and HCN results are presented here.

3. ABUNDANCES OF HCN AND C_2H_2

In order to determine the abundances of the various molecules, we first find the populations of the rotational level (N_j) from the equivalent widths, assuming optically thin lines. We can then solve for the temperature and column density. For HCN, we find that the data are best fitted by two temperature components, T_{cool}=110 K and T_{hot}=685 K, with corresponding column densities, N_{cool}=10^{16} cm^{-2} and N_{hot}=10^{16} cm^{-2}. For C_2H_2, we also find two temperature components, T_{cool}=110 K and T_{hot}=800 K. The corresponding column densities are N_{cool}=4×10^{15} cm^{-2} and N_{hot}=6×10^{15} cm^{-2}. In order to derive the abundance, we take the column density of H_2 from absorption measurements of ^{13}CO from Mitchell et al. (1990), assuming ^{12}CO/^{13}CO=85 (Wilson & Rood 1994) and ^{12}CO/H_2 = 2×10^{-4} (Lacy et al. 1994). Mitchell et al. also report two temperature components for ^{13}CO, T_{cool}=25 K and T_{cool}=176 K. (Mitchell et al. (1990) call the higher temperature T_{hot} but for comparison we will call it T_{cool}). We use the column densities of HCN, C_2H_2, and CO associated with the cool temperature (T~100 K) as they most likely come from the same region. The abundances are X(HCN)_{cool}=(1.7±0.4)×10^{-7} and X((C_2H_2)_{cool}=(6.7±0.8)×10^{-8}. We do not derive abundances for the hot component because we do not have an estimate of the H_2 column density in that component.

4. CONSTRaining CHEMICAL MODELS

With our derived abundances, we can help constrain the chemical models simulating the chemical evolution during star formation. Molecules, like HCN and C_2H_2, are good probes for the chemical models since they can be produced by grain surface chemistry and by gas-phase interactions. We compare our results with the predicted abundances from low temperature models by Rodgers & Charnley (2001) (see figures 1 and 2). The models investigate the effects of NH_3 on CH_3OH chemistry. Ice evaporation from grain mantles is included in the models for the following molecules: CO, N_2, CH_4, H_2CO, C_2H_5OH, H_2S, C_2H_6, H_2O, and CH_3OH. Destruction of CH_4 produces C_3H_2. In the two models presented here, one includes NH_3 ice evaporation while the other had no NH_3 (hereafter models 1 and 2, respectively). The core temperature is 100 K and the density is 10^7 cm^{-3}. 

![Fig. 1. Predicted abundances from assuming NH_3 evaporating from grain mantles (model 1). The horizontal lines indicate the upper and lower limits from our observations for HCN (solid) and C_2H_2 (dotted). The vertical lines indicate the range of time in which the model fits the observations. For complete model results see Rodgers & Charnley (2001).](image-url)
In figure 1, model 1 shows two different time intervals for HCN and one for C$_2$H$_2$ in which the predicted abundance is within the observed upper and lower limits of HCN (solid lines) and C$_2$H$_2$ (dashed lines) in NGC 7538 IRS 1. In addition, none of the time intervals overlap. In order for the model to fit our observations better, C$_2$H$_2$ should reach higher abundances at an earlier time in order for the time intervals for HCN and C$_2$H$_2$ to coincide. In figure 2, model 2 underpredicts the abundance of HCN we observe. This suggests that NH$_3$ ice did evaporate from grain mantles in NGC 7538 IRS 1. This model also shows two time intervals in which the predicted abundance of C$_2$H$_2$ is within our observed limits. Since the abundance of C$_2$H$_2$ depends on the initial abundance of CH$_4$, varying the amount of CH$_4$ should change the C$_2$H$_2$ abundance. Boogert et al. (1998) report the CH$_4$ ice abundance in NGC 7538 IRS 9 (a neighboring source to IRS 1) to be 8.1×10$^{-7}$, which is about twice the assumed value of the models (3.8×10$^{-7}$). Doubling the initial CH$_4$ abundance should increase the abundance of C$_2$H$_2$ but there may still be no overlap between HCN and C$_2$H$_2$. Other possible ways to increase the abundance of C$_2$H$_2$ at earlier times are for C$_2$H$_2$ to form on grains and evaporate directly from icy mantles and for C$_2$H$_2$ to be a product of evaporation (not just CH$_4$ destruction). In order to constrain the chemical models further, observations of more molecules are needed. We will be able to constrain the abundance of gas-phase CH$_4$, CH$_3$ and NH$_3$ toward NGC 7538 IRS 1 with mid-infrared observations similar to the HCN and C$_2$H$_2$ observations.

Fig. 2. Same as figure 1 but for model 2. For complete model results see Rodgers & Charnley (2001).

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REFERENCES