HCN and HNC mapping of the protostellar core Chamaeleon-MMS1*

P. P. Tennekes1,2, J. Harju1, M. Juvela1, and L. V. Tóth3,4

1 Observatory, PO Box 14, 00014 University of Helsinki, Finland
e-mail: peter.tennekes@helsinki.fi
2 Julius Institute, Utrecht University, The Netherlands
3 Department of Astronomy of the Loránd Eötvös University, Pázmány Péter sétány 1, 1117 Budapest, Hungary
4 Konkoly Observatory, PO Box 67, 1525 Budapest, Hungary

Received 18 February 2004 / Accepted 19 June 2006

ABSTRACT

Aims. The purpose of this study is to investigate the distributions of the isomeric molecules HCN and HNC and estimate their abundance ratio in the protostellar core Cha-MMS1 located in Chamaeleon I. Methods. The core was mapped in the J = 1–0 rotational lines of HCN, HNC, and HN13C. The column densities of H15CN, HN15C, H13NC and NH3 were estimated towards the centre of the core. Results. The core is well delineated in all three maps. The kinetic temperature in the core, derived from the NH3 (1, 1) and (2, 2) inversion lines, is 12.1 ± 0.1 K. The HN13C/H13CN column density ratio is between 3 and 4, i.e. similar to values found in several other cold cores. The HN13C/H13NC column density ratio is ∼7. In case no 15N fractionation occurs in HNC (as suggested by recent modelling results), the HNC/HN13C abundance ratio is in the range 30–40, which indicates a high degree of 13C fractionation in HNC. Assuming no differential 13C fractionation the HCN and HNC abundances are estimated to be ∼7 × 10−10 and ∼2 × 10−9, respectively, the former being nearly two orders of magnitude smaller than that of NH3. Using also previously determined column densities in Cha-MMS1, we can put the most commonly observed nitrogenous molecules in the following order according to their fractional abundances: γ(NH3) > γ(HCN) > γ(HNC) > γ(N2H+). Conclusions. The relationships between molecular abundances suggest that Cha-MMS1 represents an evolved chemical stage, experiencing at present the “late-time” cyanopolyyne peak. The possibility that the relatively high HNC/HCN ratio derived here is only valid for the 13C isotopic substituents cannot be excluded on the basis of the present and other available data.

Key words. ISM: individual objects: Chamaeleon-MMS1 – ISM: abundances – ISM: molecules

1. Introduction

Hydrogen cyanide, HCN, and its metastable isomer hydrogen isocyanide, HNC, are commonly used tracers of dense gas in molecular clouds. The HNC column density has been observed to be similar or larger than that of HCN in cold regions, whereas in warm GMC cores the [HNC]/[HCN] ratio is typically much smaller than unity (e.g., Goldsmith et al. 1981; Churchwell et al. 1984; Irvine & Schloerb 1984; Schilke et al. 1992; Hirota et al. 1998).

Gas-phase chemistry models, including both ion-molecule and neutral-neutral reactions, predict that the HNC/HCN abundance ratio should be close to unity in cold gas (see e.g. Herbst et al. 2000, where also previous work is reviewed). The two isomers are thought to form primarily via the dissociative recombination of HCNH+, which in turn is produced by a reaction between NH3 and C+. The recombination reaction regulates the abundance ratio despite the initial production mechanisms of the two isomers, provided that they are efficiently protonated via reactions with ions, such as H3+ and HCO+. (Churchwell et al. 1984; Brown et al. 1989). Recent theoretical studies corroborate the equal branching ratios for HCN and HNC in dissociative recombination (Hickman et al. 2005; Ishii et al. 2006).

HCN/HNC column density ratios clearly larger than unity have been reported: Churchwell et al. 1984 find that [HCN]/[HCN] may be as high as ∼10 in some dark clouds cores, and Hirota et al. (1998) find a ratio of ∼5 in L1498. The first result quoted is uncertain because it depends on the assumed degree of 13C fractionation in HNC, whereas the second value refers to HN13C and H13CN measurements. Nevertheless, both results suggest that the HCN/HNC chemistry is not fully understood. Further determinations of the relative abundances of these molecules towards cores with well-defined chemical and physical characteristics seem therefore warranted.

In this paper we report on observations of HCN and HNC and some of their isotopologues towards the dense core Cha-MMS1, which is situated near the reflection nebula Ced 110 in the central part of the Chamaeleon I dark cloud. We also derive the gas kinetic temperature in the core by using the (1, 1) and (2, 2) inversion lines of NH3. Cha-MMS1 was discovered in the 1.3 mm dust continuum by Reipurth et al. (1996), and was studied in several molecules by Kontinen et al. (2000). The object is embodied in one of the most massive “clumps” in Chamaeleon I identified in the large scale C18O survey of Haikala et al. 2005 (clump No. 3, ∼12 M⊙). Reipurth et al. (1996) suggested that Cha-MMS1 contains a Class 0 protostar. A FIR source (Ced 110 IRS10) was detected near the centre of

* Based on observations collected at the European Southern Observatory, La Silla, Chile, and at the Parkes Observatory operated by the Australia Telescope National Facility (ATNF).
the core by Lehtinen et al. (2001). However, no centimetre continuum nor near-infrared sources have been found in its neighbourhood (Lehtinen et al. 2003, and references therein), and the core therefore represents a very early stage of star formation. At the same time, its chemical composition has probably reached an advanced stage (Kontinen et al. 2000).

In Sect. 2 of this paper we describe the observations. In Sect. 3 we present the direct observational results in the form of spectra and maps, and derive the H$^{13}$CN, HN$^{15}$C, and H$^{15}$NC column densities. By combining our results with previous observations we derive in Sect. 4 the fractional abundances of several nitrogen containing molecules in Cha-MMS1, and discuss briefly the chemical state of the core. In Sect. 5 we summarize our results.

2. Observations

The $J = 1$–0 transitions of HCN, H$^{13}$CN, HNC, HN$^{15}$C and H$^{15}$NC at about 90 GHz were observed with the Swedish-ESO-Submillimeter-Telescope, SEST, located on La Silla in Chile. The observations took place in December 1990. A 3 mm dual polarization Schottky receiver was used in the frequency switching mode. The backend was a 2000 channel acousto optical spectrometer with an 86 MHz bandwidth. The velocity resolution with this configuration is about 0.15 km s$^{-1}$ at 90 GHz. Typical values for the single sideband system temperatures ranged from 350 K to 450 K. The half-power beam width of the SEST is about 55$''$ at the frequencies observed. Calibration was achieved by the chopper wheel method. Pointing was checked every 2–3 h towards a nearby SiO maser source. We estimate the pointing accuracy to have been better than 5$''$ during the observations. The focusing was done using a strong SiO maser.

The (1, 1) and (2, 2) transition lines of NH$_3$ at about 23 GHz were observed with the Parkes 64-m radio telescope of the Australian Telescope National Facility (ATNF) in April 2000. The FWHM of the telescope is about 80$''$ at this frequency. The two lines were observed simultaneously in two orthogonal polarizations in the frequency switching mode, using the 8192 channel autocorrelator. The calibration was checked by observing standard extragalactic calibration sources at different elevations, and by comparing the ammonia line intensities towards some strong galactic sources also visible from the Effelsberg 100-m telescope, taking the different beam sizes into account.

The electric dipole moments of the molecules and the line frequencies used in this paper are listed in Table 1. Except for the frequencies of the HNC and HN$^{15}$C hyperfine transitions adopted from Frerking et al. (1979), the parameters are obtained from the Jet Propulsion Laboratory molecular spectroscopy database (http://spec.jpl.nasa.gov), where also the original references are given. The hyperfine components of NH$_3$ are not listed. The frequencies given represent the centre frequencies of 18 and 21 individual components of the (1, 1) and (2, 2) lines, respectively, distributed over a range of about 4 MHz.

3. Maps and column densities

3.1. Maps

The integrated HCN($J = 1$–0), HNC($J = 1$–0) and HN$^{15}$C($J = 1$–0) intensity maps of Cha-MMS1 are presented as contour plots in Figs. 1–3, respectively. The values of the contour levels are given in the bottom right of each figure. The position of the 1.3 mm dust emission maximum, Cha-MMS1a

![Fig. 1. Integrated HCN($J = 1$–0) line intensity map. The integration is performed over all three hyperfine structure components ($F = 1$–1, $F = 2$–1 and $F = 0$–0). The map is centered on the position of Cha-MMS1a (denoted by a cross), $\alpha_{2000} = 11^h06^m31^s7$, $\delta_{2000} = -77^\circ23'32''$, and the offsets are in arcseconds. The contour levels indicated with ten shades of grey and their values can be read in the bottom right corner. The intensity unit is K km s$^{-1}$. The positions used for the column density estimates (–20$''$, 20$''$, and T$_{ex}$ estimates (see Table 2) are marked with a black dot and open circles, respectively. The grid spacing used in the map is 30$''$.](http://spec.jpl.nasa.gov)

Fig. 1. Integrated HCN($J = 1$–0) line intensity map. The integration is performed over all three hyperfine structure components ($F = 1$–1, $F = 2$–1 and $F = 0$–0). The map is centered on the position of Cha-MMS1a (denoted by a cross), $\alpha_{2000} = 11^h06^m31^s7$, $\delta_{2000} = -77^\circ23'32''$, and the offsets are in arcseconds. The contour levels indicated with ten shades of grey and their values can be read in the bottom right corner. The intensity unit is K km s$^{-1}$. The positions used for the column density estimates (–20$''$, 20$''$, and T$_{ex}$ estimates (see Table 2) are marked with a black dot and open circles, respectively. The grid spacing used in the map is 30$''$. (Reipurth et al. 1996), is chosen as the map centre. The coordinates of Cha-MMS1a are $\alpha_{2000} = 11^h06^m31^s7$, $\delta_{2000} = -77^\circ23'32''$. The grid spacing used in the maps was 30$''$. Also indicated in these figures are the location of the HN$^{15}$C maximum ($\Delta x, \Delta y = -20''$, 20$''$, black dot), and the positions used for the HCN($J = 1$–0) excitation temperature estimates in the LTE method (open circles, see below). Long integrations towards the HN$^{15}$C maximum were made besides HNC and HCN also in the $J = 1$–0 lines of H$^{13}$CN and H$^{15}$NC. The five spectra obtained towards this position are shown in Fig. 5.

The three maps have differences, but they all show a relatively compact distribution peaking near Cha-MMS1a. The HN$^{15}$C emission with the lowest opacity reflects probably best the distribution dense gas.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>$\mu$ (Debye)</th>
<th>Transition</th>
<th>$\nu$ (MHz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCN</td>
<td>2.98</td>
<td>$J = 1$–0</td>
<td>88630.416</td>
</tr>
<tr>
<td>H$^{13}$CN</td>
<td>2.98</td>
<td>$J = 1$–0</td>
<td>88631.847</td>
</tr>
<tr>
<td>HNC$^b$</td>
<td>3.05</td>
<td>$J = 1$–0</td>
<td>86340.184</td>
</tr>
<tr>
<td>HN$^{15}$C$^b$</td>
<td>2.70</td>
<td>$J = 1$–0</td>
<td>87090.942</td>
</tr>
<tr>
<td>H$^{15}$NC</td>
<td>2.70</td>
<td>$J = 1$–0</td>
<td>88865.715</td>
</tr>
<tr>
<td>NH$_3$</td>
<td>1.48</td>
<td>(J, K) = (1, 1)</td>
<td>23694.495</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(J, K) = (2, 2)</td>
<td>23722.633</td>
</tr>
</tbody>
</table>

$^b$ Frequencies from Frerking et al. (1979).

Table 1. The permanent dipole moments of the molecules and the rest frequencies of the observed transitions.
beams brightness temperature, $T_{MB}$ and assuming uniform beam filling. The fit is possible towards the four positions listed in Table 2. In other positions with strong signal, like the HN$^{13}$C maximum ($-20^\prime\prime$, $+20^\prime\prime$), or the HCN maximum at ($-20^\prime\prime$, $-20^\prime\prime$) the HCN spectra show “anomalous” hyperfine ratios. The smallest error is obtained towards the offset ($40^\prime\prime$, $20^\prime\prime$) near the edge of the core with $T_{ex}$ $\approx 5$ K. This HCN spectrum is shown in Fig. 4. The other positions listed in Table 2 give similar values of $T_{ex}$ but the errors are larger. Also HCN column density estimates using the values for $\tau$ and $\Delta \delta$ obtained from hyperfine fits are given in Table 2.

The HCN, H$^{13}$CN, HN$^{13}$C and H$^{15}$NC column densities towards the HN$^{13}$C maximum, ($-20^\prime\prime$, $+20^\prime\prime$), have been derived from the integrated $J = 1–0$ line intensities and the $T_{ex}$ estimate derived from the HCN spectrum shown in Fig. 4. For HCN the weakest of the well separated hyperfine components, $F = 0–1$, with the least optical thickness has been used in this calculation by taking its relative strength into account. For HNC we did not attempt to do a similar estimate because the hyperfine components overlap and are very likely optically thick. Overlapping hyperfine components hamper also the HN$^{13}$C column density estimate, but their optical thicknesses should be lower than those of HNC by an order of magnitude or more. H$^{15}$NC has no hyperfine structure due to zero nuclear spin of $^{15}$N. The results of these calculations are presented in Table 3.

The following column density ratios can be derived from Table 3:

\[
\frac{N[\text{HN}^{13}\text{C}]}{N[\text{H}^{13}\text{CN}]} = 3.3 \pm 0.4
\]
\[
\frac{N[\text{HN}^{13}\text{C}]}{N[\text{H}^{15}\text{NC}]} = 7.4 \pm 1.8
\]
\[
\frac{N[\text{HCN}]}{N[\text{H}^{15}\text{CN}]} = 17 \pm 2.
\]

3.2. LTE column densities

The column densities of the molecules have been determined with two different methods. First, we have derived the optical thicknesses of the lines and the total column densities of the molecules by assuming local thermodynamic equilibrium, LTE. The LTE method involves idealistic assumptions about the homogeneity of the cloud. In the next subsection we present modelling of the observed spectra with a Monte-Carlo program using a realistic density distribution.

In the following, we assume that the rotational levels of all five isotopologues are characterized by the same $T_{ex}$, and that the core is homogeneous, i.e. the same $T_{ex}$ is valid for different locations. The former assumption can be justified by the similar dipole moments and rotational constants of the molecules. The second assumption infers that the excitation conditions remain constant along the line of sight and also between close-lying positions in the core.

The excitation temperature is derived by fitting first the optical thickness, $\tau$, to the observed hyperfine ratios of a “clean” HCN spectrum, i.e. a spectrum with a good $S/N$ and hyperfine intensity ratios in accordance with the LTE assumption. The $T_{ex}$ is solved from the antenna equation using observed peak main

---

Table 2. Estimates of the HCN($J = 1–0$) excitation temperatures and the HCN column densities towards selected positions using hyperfine structure fits. The coordinates are offsets from the position of Cha-MMS1.

<table>
<thead>
<tr>
<th>$\Delta \alpha$</th>
<th>$\Delta \delta$</th>
<th>$T_{ex}$</th>
<th>$\tau_{\text{main}}$</th>
<th>$\Delta v$</th>
<th>$N$(HCN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>40</td>
<td>20</td>
<td>5.2$^{+1.0}_{-0.8}$</td>
<td>1.1 $\pm$ 0.3</td>
<td>0.49 $\pm$ 0.30</td>
<td>4.7 $\times 10^{12}$</td>
</tr>
<tr>
<td>-50</td>
<td>-10</td>
<td>5.3$^{+1.0}_{-0.8}$</td>
<td>0.9 $\pm$ 0.4</td>
<td>0.58 $\pm$ 0.07</td>
<td>4.6 $\times 10^{12}$</td>
</tr>
<tr>
<td>10</td>
<td>-10</td>
<td>5.6$^{+1.0}_{-0.8}$</td>
<td>0.7 $\pm$ 0.3</td>
<td>0.56 $\pm$ 0.08</td>
<td>3.8 $\times 10^{12}$</td>
</tr>
<tr>
<td>70</td>
<td>20</td>
<td>5.4$^{+1.0}_{-0.8}$</td>
<td>0.5 $\pm$ 0.3</td>
<td>0.49 $\pm$ 0.04</td>
<td>2.0 $\times 10^{12}$</td>
</tr>
</tbody>
</table>

---

Fig. 2. Integrated HNC($J = 1–0$) line intensity map. The coordinates and markers as in Fig. 1.

Fig. 3. Integrated HN$^{13}$C($J = 1–0$) line intensity map. The coordinates and markers as in Fig. 1.

Fig. 4. The HCN($J = 1–0$) spectrum towards offset ($40^\prime\prime$, $20^\prime\prime$) used to determine the $T_{ex}$. The dotted line represents a fit to the hyperfine structure.
The ammonia column density, $N(\text{NH}_3)$, and the gas kinetic temperature, $T_{\text{kin}}$, were derived using hyperfine fits to the NH$_3$ inversion lines arising from the rotational states $(J, K) = (1, 1)$ and (2, 2). The spectra are shown in Fig. 6. The standard method of analysis described e.g. in Ho et al. (1979) gave the following values: $N(\text{NH}_3) = (1.4 \pm 0.2) \times 10^{19} \text{cm}^{-2}$ and $T_{\text{kin}} = 12.1 \pm 0.1 \text{K}$. Here it is assumed that the relative populations of all metastable levels of both ortho- and para-NH$_3$ are determined by thermal equilibrium at the (2, 2)/(1, 1) rotational temperature.

The HCN/H$^{13}$CN column density ratio, 17, is much lower than the isotopic $^{12}$C/$^{13}$C ratios usually observed in nearby molecular clouds or local diffuse ISM (~60, e.g. Lucas & Liszt 1998; the terrestrial value is 89), and is likely to be affected by absorption or scattering in the less dense envelope of the core. Radiative transfer effects are even more obvious in the case of the HNC/H$^{13}$C intensity ratio, 1.7. We do not use therefore the values derived from the HCN and HNC spectra towards the core centre. The HCN and HNC spectra are discussed at the end of Sect. 4.

The ammonia column density, $N(\text{NH}_3)$, and the gas kinetic temperature, $T_{\text{kin}}$, were derived using hyperfine fits to the NH$_3$ inversion lines arising from the rotational states $(J, K) = (1, 1)$ and (2, 2). The spectra are shown in Fig. 6. The standard method of analysis described e.g. in Ho et al. (1979) gave the following values: $N(\text{NH}_3) = (1.4 \pm 0.2) \times 10^{19} \text{cm}^{-2}$ and $T_{\text{kin}} = 12.1 \pm 0.1 \text{K}$. Here it is assumed that the relative populations of all metastable levels of both ortho- and para-NH$_3$ are determined by thermal equilibrium at the (2, 2)/(1, 1) rotational temperature.
densities. The H$^{13}$CN and HN$^{13}$C column densities (fractional abundances multiplied by N(H$_2$)) are indeed about two times larger than those presented in the previous section: N(H$^{13}$CN) = 1.1 × 10$^{12}$ cm$^{-2}$, N(HN$^{13}$C) = 4.0 × 10$^{12}$ cm$^{-2}$. These results suggest, however, that the LTE method does not lead to crude underestimates of the column densities of rarer isotopologues of HCN and HNC.

4. Discussion

4.1. Fractional abundances

The present observations and the LTE method give reasonable estimates for the column densities of H$^{13}$CN, HN$^{13}$C, H$^{15}$NC and NH$_3$ near the centre of Cha-MMS1. Unfortunately the column densities of the common isotopologues of HCN and HNC cannot be directly determined because of the large optical thickness of their lines. H$^{15}$NC is, however, likely to be useful for this purpose. According to the modelling results of Terzieva & Herbst (2000) hardly any $^{15}$N fractionation occurs in dense interstellar clouds. The isotopic $^{14}$N/$^{15}$N ratio lies probably between 240 and 280, i.e. the values derived for the local ISM (Lucas & Liszt 1998), and the terrestrial atmosphere. Using this range and the H$^{15}$NC column density listed in Table 3 we obtain N(NHC) ~ 6 × 10$^{13}$ cm$^{-2}$. Assuming that no differential $^{13}$C fractionation occurs, we get from the HN$^{13}$C/H$^{13}$CN ratio derived above that N(HCN) ~ 2 × 10$^{13}$ cm$^{-2}$.

The dust continuum emission provides the most reliable estimate for the H$_2$ column density, Reipurth et al. (1996) measured towards Cha-MMS1 a total 1.3 mm flux density of 950 mJy from a nearly circular region with a diameter of about 50” (bordered by the lowest white contour in Fig. 8). As this size is similar to the SEST beamsize at 3 mm, the resulting average intensity and average N(H$_2$) can be compared with the column densities derived here. The dust temperature as derived from ISO observations by Lehtinen et al. (2001) is 20 K (Cha-MMS1  = Ced 110 IRS10). Due to crowding of sources in this region and the limited angular resolution of ISOPHOT, this value may be affected by emission from other sources in its vicinity (see Fig. 2 of Lehtinen et al. 2001). Therefore, value 20 K is probably an upper limit. A lower limit is provided by the gas kinetic temperature ~12 K derived above. By using the two temperatures and Eq. (1) of Motte et al. 1998 with $\kappa_{3.3mm} = 0.01$ g cm$^{-2}$ appropriate for circumstellar envelopes around Class 0 and Class I protostars, we get N(H$_2$) = 1.9 – 4.0 × 10$^{22}$ cm$^{-2}$. The corresponding mass range of the dust core obtained from the total flux given by Reipurth et al. (1996) is 0.4–0.8 $M_\odot$.

The median of the reasonable H$_2$ column density range, N(H$_2$) = 3.0×10$^{22}$ cm$^{-2}$, yields the following values for the fractional HCN, HNC and NH$_3$ abundances: $\chi$(HCN) ~ 7×10$^{-10}$, $\chi$(HNC) ~ 2×10$^{-9}$, and $\chi$(NH$_3$) ~ 5×10$^{-8}$. The column densities of two other nitrogen bearing molecules towards Cha-MMS1 were determined by Kontinen et al. (2000): N(HC$_3$N) = (4.5 ± 0.8) × 10$^{14}$ cm$^{-2}$, and N(N$_2$H$^+$) = (1.4 ± 0.1) × 10$^{13}$ cm$^{-2}$, which imply the fractional abundances $\chi$(HC$_3$N) ~ 2×10$^{-8}$ and $\chi$(N$_2$H$^+$) ~ 5×10$^{-10}$.

We find that both HCN and N$_2$H$^+$ are about 100 times less abundant than NH$_3$, whereas HC$_3$N is almost equally abundant as NH$_3$. The derived N$_2$H$^+$/NH$_3$ column density ratio is in accordance with previous observational results (e.g., Tafalla et al. 2002; Hotzel et al. 2004). The abundance ratios are discussed in Sect. 4.3. The C$^{18}$O column density in this direction is (1.5 ± 0.1) × 10$^{15}$ cm$^{-2}$ (Haikala et al. 2005). The fractional C$^{18}$O abundance is therefore 5 × 10$^{-8}$ or smaller. This upper limit falls below the “standard” value ~2×10$^{-7}$ of Frerking et al. (1982), and suggests CO depletion.

4.2. Comparison with previous maps

The superposition of our HN$^{13}$C map, the HC$_3$N map of Kontinen et al. (2000), and the 1.3 mm dust continuum map of Reipurth et al. (1996) is shown in Fig. 8. The two molecular line maps have similar angular resolution (about 60”) whereas the resolution of the continuum map is 22”. Both molecules show rather compact, symmetric distributions around the dust continuum source, HC$_3$N agreeing slightly better with dust than HN$^{13}$C. Nevertheless, the dense core can be clearly localized with the aid of HN$^{13}$C as well as HCN and HNC. This is in contrast with SO, which shows two peaks on both sides of the core (Kontinen et al. 2000), and with C$^{18}$O which has a flat distribution in this region (Haikala et al. 2005), probably due to depletion. This result agrees with the results of Hirota et al. (2003) and Nikolić et al. (2003), which suggest that HCN and HNC are less affected by the accretion onto dust grains than some other commonly observed molecules.
4.3. The chemical state of Cha-MMS1 and the HNC/HCN abundance ratio

CO depletion and a large NH₃ abundance point towards an advanced stage of chemical evolution. The NH₃/N₂H⁺ abundance ratio ∼100 agrees with model predictions for dense cores, but does not put severe constraints on the evolutionary stage (Aikawa et al. 2005). According to the observational results of Hotzel et al. (2004) this value should be more appropriate for a star-forming than for a prestellar core. However, due to slightly different beam sizes the accuracy of the present determination of the NH₃/N₂H⁺ ratio is not sufficient for distinguishing between the two cases. Kontinen et al. (2000) suggested that the large HC₃N abundance observed in Cha-MMS1 indicates either a so called “late-time” cyanopolyene peak associated with a large degree of depletion according to the model of Ruffle et al. (1997), or carbon chemistry revival by the influence of a newly born, so far undetected star. A large ammonia abundance can be understood in either case. Ammonia production benefits from the diminishing of CO and can resist depletion longer (e.g. Nejad & Wagenblast 1999). On the other hand, ammonia can possibly form on grain surfaces and be released into the gas phase in shocks (Nejad et al. 1990; Aikawa et al. 2003). As there is no clear evidence for star-cloud interaction in Cha-MMS1, the alternative that the core – or the portion traced by the present observations – represents matured pre-stellar chemistry is more likely.

The behaviour of HCN and HNC under these circumstances is not clear. The main gas-phase production pathway via the HCNH⁺, which is effective especially at later stages, requires NH₃ and C⁺ ions. The latter can be produced also in deep interiors of molecular clouds via the cosmic ray ionization of C atoms, or in the reaction between CO and the He⁺ ion, the latter mechanism becoming less important with CO depletion (Nejad & Wagenblast 1999). To our knowledge the production of HCN and HNC on interstellar grains and their desorption has not been studied.

The models of Aikawa et al. (2005) for collapsing prestellar cores include the gas-phase production of HCN, HC₃N, NH₃, and N₂H⁺. Although the order of fractional abundances, \( \chi(\text{NH}_3) > \chi(\text{HCN}) > \chi(\text{N}_2\text{H}^+) \), and the \( \text{NH}_3/\text{N}_2\text{H}^+ \) abundance ratio derived from the present observations are well reproduced by this model, especially for the gravity-to-pressure ratio \( \alpha = 4 \) implying a rapid collapse and less marked depletion, the predicted HCN column densities relative to \( \text{NH}_3 \) and \( \text{N}_2\text{H}^+ \) are an order of magnitude larger than those observed here. A likely explanation is that HCN is not as centrally peaked than \( \text{NH}_3 \) and \( \text{N}_2\text{H}^+ \) (see Fig. 3 of Aikawa et al. 2005). Therefore a large fraction of HCN resides in on the outskirts of the core, where the density is not sufficient for the collisional excitation of its rotational levels, and the molecule is not seen in emission (see also the next section). Another discrepancy is that the model predicts much lower \( \text{HC}_3\text{N} \) column densities relative to ammonia than observed here. The agreement is slightly better for the model with overwhelmingly large gravitational potential \( (\alpha = 4) \) than for the slowly evolving model near hydrostatic equilibrium \( (\alpha = 1.1) \) which results in very high degrees of depletion. Unfortunately the spectral resolution and S/N ratio of the present data do not allow to study the core kinematics.

As compared to the “principal” N-bearing molecules \( \text{NH}_3 \) and \( \text{HC}_3\text{N} \), HCN and HNC have clearly smaller abundances, and probably only marginal importance to the nitrogen chemistry in the core. Nevertheless, the fact that HNC/HCN abundance ratio is once again observed to be larger than unity remains a problem. Talbi & Herbst (1998) have shown that even though the reaction between \( \text{NH}_3 \) and \( \text{C}^+ \) may produce a significant amount of the metastable ion \( \text{H}_2\text{NC}^- \), which should produce only HNC in electron recombination, the energy released in the former reaction leads to the efficient transformation into the linear isomer \( \text{HCN}^+ \) before the recombination. Likewise, according to Herbst et al. (2000), the vibrational energies of HNC and HCN formed in the neutral reactions \( \text{C} + \text{NH}_2 \rightarrow \text{HNC} + \text{H} \), and \( \text{N} + \text{CH}_2 \rightarrow \text{HCN} + \text{H} \) are sufficient to overcome the isomerization barrier, and roughly equal amounts of both isomers are produced.

The theoretical results presented in Talbi & Herbst (1998) and Herbst et al. (2000), and more recently by Hickman et al. (2005) and Ishii et al. (2006), show quite convincingly that the HNC/HCN abundance ratio resulting from the chemical reactions known to be effective in dark clouds cannot differ significantly from unity. There are two possible explanations for the discrepancy between the observations and theoretical predictions. Either so far unknown processes favour HNC in the cost of HCN, or then the derived HN\(^{13\text{C}}\)/HC\(^{13\text{C}}\) column density ratio is affected by different degrees of \(^{13\text{C}}\) fractionation in HNC and HCN. In the latter case \(^{13\text{C}}\) would about three times more enhanced in HNC than in HCN in Cha-MMS1.

The HN\(^{13\text{C}}\)/HC\(^{13\text{C}}\) column density ratio 7.4 together with the above quoted range of \( ^{14\text{N}}/^{13\text{N}} \) ratios imply that the HNC/HCN ratio in Cha-MMS1 is in the range 30–40. As the typical \( ^{12}\text{C}^{13}\text{C} \) isotopic ratio observed in nearby clouds is 60 (Lucas & Liszt 1998) the result suggests considerable \(^{13}\text{C} \) enhancement in HNC. An HC\(^{15}\text{N} \) column density determination would readily show, according to the results of Terzieva & Herbst (2000), whether also HCN has a similar degree of \(^{13}\text{C} \) fractionation.

In the only survey so far including all these isotopologues, Irvine & Schloerb (1984) found very similar HN\(^{13}\text{C}/\text{HC}^{13}\text{N} \) and H\(^{15}\text{N}/\text{HC}^{15}\text{N} \) intensity ratios (~1.5) towards TMC-1. This observation suggests similar \(^{13}\text{C} \) fractionation in HCN and HNC. On the other hand, this result might not be universally valid. Ikeda et al. 2002 determined substantially different H\(^{13}\text{CN}/\text{HC}^{13}\text{N} \) column density ratios in the starless dark cloud cores L1521E, TMC-1, and L1498, listed in order of increasing H\(^{13}\text{CN}/\text{HC}^{15}\text{N} \) ratio (2.6, 6.7, and 11). The H\(^{13}\text{C}/\text{HC}^{15}\text{N} \) column density ratios in these sources increase in the same order (0.54, 1.7, and 4.5; Hirota et al. 1998). The values obtained towards TMC-1 are consistent with those of Irvine & Schloerb (1984), but the results towards L1521E and L1498 suggest (if the possibility of \(^{15}\text{N} \) fractionation is neglected) that the \(^{13}\text{C} \) fractionation in HCN changes from source to source, and perhaps the variation in the case of HNC is even larger. L1521E is suggested to represent youthfull chemistry with little molecular depletion (Hirota et al. 2002; Tafalla & Santiago 2004) whereas the position observed in L1498 is known to be heavily depleted (Willacy et al. 1998). Thus, it appears that the freeze-out of molecules favour the \(^{13}\text{C} \) substituted isotopologues of HCN and HNC. It seems warranted to further examine the possibility of their differential \(^{13}\text{C} \) fractionation, although no theoretical arguments are in favour of different gas-phase production and destruction rates of HN\(^{13}\text{C} \) and H\(^{15}\text{CN} \), or their different freezing rates. As no empirical data on this matter is currently available, the best way to proceed is probably to perform similar observations as those done by Irvine & Schloerb (1984) towards a sample of cores with different degree of depletion.
4.4. Scattering envelope

The low HCN/H$^{13}$CN and HNC/HN$^{13}$C intensity ratios and the line shapes in the HCN and HNC spectra observed towards the centre of the core require some attention. The intensity ratios suggest very large optical thicknesses for the HCN and HNC lines, and consequently, deep self-absorption features and “anomalous” hyperfine component ratios are to be expected. A clear absorption feature can be seen in the simulated HCN spectrum shown in Fig. 7, though a very low $^{12}\text{C}/^{13}\text{C}$ isotopic ratio is used in order to avoid inverted hyperfine ratios, which would disagree with the observations. The observed HCN and HNC spectra are, however, free from absorption dips, and they look more like weakened “normal” lines.

This weakening can probably be attributed to less dense gas surrounding the core. The HCN and HNC molecules in the envelope, where the density lies below the critical density of their $J = 1–0$ lines, function as scatterers of line emission arising from the core, i.e. they re-emit absorbed photons (into random directions) before colliding with other particles. The effect is less marked for $^{13}$C isotopologues, and the resulting HCN/H$^{13}$CN and HNC/HN$^{13}$C intensity ratios decrease, without HCN and HNC losing their characteristic line patterns. This model was used by Cernicharo et al. (1984) to explain HCN hyperfine anomalies, particularly the intensity of the intrinsically weakest component $F = 0–1$ observed towards dark clouds. The HCN spectrum shown in Fig. 5 conforms with this explanation: the $F = 0–1$ component on the left is stronger than $F = 1–1$ component on the right, although the optically thin LTE ratio is 1:3.

The scattering of HCN emission from the core has probably lead to underestimation of the excitation temperature and column densities derived using the LTE method in Sect. 3.2. For example, an increase of 1 K in $T_{\text{ex}}$, would mean 30% larger column densities, and 3 K would correspond to a 100% increase in the column density. Quantitative estimates of the scattering effects are, however, beyond the scope of this paper, and here we merely note this additional source of error.

5. Summary

A mapping in the $J = 1–0$ rotational lines of HCN, HNC and HN$^{13}$C shows a compact structure in the direction of the protostellar core Cha-MMS1. The HN$^{13}$C map agrees best with the results of Terzieva & Herbst (2000), we estimate that the [HNC]/[HN$^{13}$C] ratio is between 30 and 40. As there is no HCN/HNC ratio determination available towards this source, the possibility that differential $^{13}$C fractionation causes the observed relatively high [HN$^{13}$C]/[H$^{13}$CN] ratio cannot be entirely ruled out.

The HCN and HNC lines are not much brighter than those of the $^{13}$C substituted species, but still show no absorption dips. These lines are probably weakened by scattering by HCN and HNC molecules residing in less dense gas around the core. As suggested by Cernicharo et al. (1984), the same process can cause the so called HCN hyperfine anomalies in dark clouds, and is probably also responsible for the “too” bright $F = 0–1$ component of HCN ($J = 1–0$) towards the centre of Cha-MMS1. The evidence for scattering and the relatively low HCN and HNC column densities derived towards the core suggest that the HCN and HNC distributions are less centrally peaked than those of NH$_3$ and N$_2$H$^+$, which would be in agreement with the model of Aikawa et al. (2005) for a rapidly collapsing core.

Acknowledgements. We thank Wolf Geppert for useful discussions and the anonymous referee for valuable comments on the manuscript. This project was supported by the Academy of Finland, grant Nos. 173727 and 174854, P.T. thanks the ERASMUS programme of the European Union, Olga Koningsfonds, Utrecht, and Kapteyn Fonds, Groningen.

References


The following column density ratios are found: HN$^{13}$C/H$^{13}$CN $\sim$ 3–4 and HN$^{13}$C/H$^{15}$NC $\sim$ 7. Using the latter ratio, and the assumption based on the modelling results of Terzieva & Herbst (2000), we estimate that the [HNC]/[HN$^{13}$C] ratio is between 30 and 40. As there is no HCN/HNC ratio determination available towards this source, the possibility that differential $^{13}$C fractionation causes the observed relatively high [HN$^{13}$C]/[H$^{13}$CN] ratio cannot be entirely ruled out.