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Shedding light on the formation of the pre-biotic molecule formamide with ASAI

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ABSTRACT

Formamide (NH₂CHO) has been proposed as a pre-biotic precursor with a key role in the emergence of life on Earth. While this molecule has been observed in space, most of its detections correspond to high-mass star-forming regions. Motivated by this lack of investigation in the low-mass regime, we searched for formamide, as well as isocyanic acid (HNCO), in 10 low- and intermediate-mass pre-stellar and protostellar objects. The present work is part of the IRAM Large Programme ASAI (Astrochemical Surveys At IRAM), which makes use of unbiased broad-band spectral surveys at millimetre wavelengths. We detected HNCO in all the sources and NH₂CHO in five of them. We derived their abundances and analysed them together with those reported in the literature for high-mass sources. For those sources with formamide detection, we found a tight and almost linear correlation between HNCO and NH_2CHO abundances, with their ratio being roughly constant – between 3 and 10 – across 6 orders of magnitude in luminosity. This suggests the two species are chemically related. The sources without formamide detection, which are also the coldest and devoid of hot corinos, fall well off the correlation, displaying a much larger amount of HNCO relative to NH₂CHO. Our results suggest that, while HNCO can be formed in the gas-phase during the cold stages of star formation, NH2CHO forms most efficiently on the mantles of dust grains at these temperatures, where it remains frozen until the temperature rises enough to sublimate the icy grain mantles. We propose hydrogenation of HNCO as a likely formation route leading to NH₂CHO.

Key words: astrochemistry – methods: observational – stars: formation – ISM: abundances – ISM: molecules.

1 INTRODUCTION

One of the major questions regarding the origin of life on Earth is whether the original chemical mechanism that led from sim-

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ple molecules to life was connected to metabolism or to genetics, both intimately linked in living beings. Formamide (NH₂CHO) contains the four most important elements for biological systems: C, H, O, and N, and it has recently been proposed as a prebiotic precursor of both metabolic and genetic material, suggesting a common chemical origin for the two mechanisms (Saladino et al. 2012).

Table 1. Source sample and their properties.

Source	R.A.(J2000)	Dec.(J2000)	$V_{\rm lsr}$ (km s ⁻¹)	<i>d</i> (pc)	M (M _{\odot})	L_{bol} (L _O)	Type*	References
ASAI								
L1544	05:04:17.21	+25:10:42.8	+7.3	140	2.7	1.0	PSC	1,2,3
TMC1	04:41:41.90	+25:41:27.1	+6.0	140	21	_	PSC	1,4
B1	03:33:20.80	+31:07:34.0	+6.5	200	1.9	1.9	Class 0	5,6
L1527	04:39:53.89	+26:03:11.0	+5.9	140	0.9	1.9	Class 0, WCCC	1,7,8
L1157-mm	20:39:06.30	+68:02:15.8	+2.6	325	1.5	4.7	Class 0	7,8
IRAS 4A	03:29:10.42	+31:13:32.2	+7.2	235	5.6	9.1	Class 0, HC	7,8
SVS13A	03:29:03.73	+31:16:03.8	+6.0	235	0.34	21	Class 0/1	9,10
OMC-2 FIR 4	05:35:26.97	-05:09:54.5	+11.4	420	30	100	IM proto-cluster	11,12
Cep E	23:03:12.80	+61:42:26.0	- 10.9	730	35	100	IM protostar	13
TIMASSS								
I16293	16:32:22.6	-24:28:33	+4.0	120	3	22	Class 0, HC	14,15

Notes. *PSC: pre-stellar core; HC: hot corino; WCCC: warm carbon-chain chemistry; IM: intermediate-mass. References: ¹Elias (1978), ²Evans et al. (2001), ³Shirley et al. (2000), ⁴Tóth et al. (2004), ⁵Hirano et al. (1999), ⁶Marcelino et al. (2005), ⁷Kristensen et al. (2012), ⁸Karska et al. (2013), ⁹Hirota et al. (2008), ¹⁰Chen, Launhardt & Henning (2009), ¹¹Crimier et al. (2009), ¹²Furlan et al. (2014), ¹³Crimier et al. (2010a), ¹⁴Loinard et al. (2008), ¹⁵Correia, Griffin & Saraceno (2004).

Formamide was detected for the first time in space by Rubin et al. (1971) towards Sgr B2 and later in Orion KL. However, dedicated studies of NH₂CHO in molecular clouds have started only very recently, as its potential as a key prebiotic molecule has become more evident. These studies present observations of formamide in a number of massive hot molecular cores (Bisschop et al. 2007; Adande, Woolf & Ziurys 2011), the low-mass protostellar object IRAS 16293–2422 (Kahane et al. 2013), and the outflow shock regions L1157-B1 and B2 (Yamaguchi et al. 2012; Mendoza et al. 2014). Its detection in comet Hale-Bopp has also been reported (Bockelée-Morvan et al. 2000). Formamide is therefore present in a variety of star-forming environments, as well as on a comet of the Solar system. Whether this implies an exogenous delivery on to a young Earth in the past is a suggestive possibility that needs more evidence to be claimed.

Establishing the formation route(s) of formamide in space remains a challenge. Different chemical pathways have been proposed, both in the gas-phase (e.g. Redondo, Barrientos & Largo 2014) and on grain surfaces (e.g. Raunier et al. 2004; Jones et al. 2011). The present work represents an effort to try to understand the dominant mechanisms that lead to the formation of formamide in the interstellar medium. In particular, it seeks to investigate the possible chemical connection between NH2CHO and HNCO, which was proposed by Mendoza et al. (2014). To this aim, we have performed a homogeneous search of NH2CHO and HNCO in a representative sample of 10 star-forming regions (SFRs) of low- to intermediatemass type, since most of the formamide detections so far reported concentrate on high-mass SFRs. This is the first systematic study conducted within the context of the IRAM Large Program ASAI (Astrochemical Surveys At IRAM; P.I.s: B. Lefloch, R. Bachiller), which is dedicated to millimetre astrochemical surveys of low-mass SFRs with the IRAM 30-m telescope.

The source sample and the observations are described in Sections 2 and 3, respectively. Section 4 presents the spectra and describes the analysis carried out to obtain the abundances of NH₂CHO and HNCO. Section 5 compares the derived abundances with those found in the literature for other SFRs, and discusses the formation mechanisms that are favoured by our results. Our conclusions are summarized in Section 6.

2 SOURCE SAMPLE

Our source sample consists of 10 well-known pre-stellar and protostellar objects representing different masses and evolutionary states, thus providing a complete view of the various types of objects encountered along the first phases of star formation. Their basic properties are listed in Table 1. All of them belong to the ASAI source sample except one: the Class 0 protobinary IRAS 16293–2422 (hereafter I16293), whose millimetre spectral survey, TIMASSS (The IRAS16293–2422 Millimeter And Submillimeter Spectral Survey), was published by Caux et al. (2011a). A dedicated study of Complex Organic Molecules (COMs) in this source, including NH₂CHO, was recently carried out by Jaber et al. (2014).

3 OBSERVATIONS AND DATA REDUCTION

The data presented in this work were acquired with the IRAM 30-m telescope near Pico Veleta (Spain) and consist of unbiased spectral surveys at millimetre wavelengths. These are part of the Large Programme ASAI, whose observations and data reduction procedures will be presented in detail in an article by Lefloch & Bachiller (in preparation). Briefly, we gathered the spectral data in several observing runs between 2011 and 2014 using the EMIR receivers at 3 mm (80-116 GHz), 2 mm (129-173 GHz), and 1.3 mm (200-276 GHz). The main beam sizes for each molecular line analysed are listed in Tables B1 and B2. The three bands were covered for most of the sources. For Cep E, additional observations were carried out at 0.9-mm (E330 receiver), while just a few frequencies were covered at 2 mm. The Fourier Transform Spectrometer (FTS) units were connected to the receivers, providing a spectral resolution of 195 kHz, except in the case of L1544, for which we used the FTS50 spectrometer, with a resolution of 50 kHz, to resolve the narrow lines ($\Delta V \sim 0.5 \text{ km s}^{-1}$) that characterize this region. The observations were performed in wobbler switching mode with a throw of 180 arcsec.

The data were reduced with the package CLASS90 of the GILDAS software collection.¹ Through comparison of line intensities among

¹ http://www.iram.fr/IRAMFR/GILDAS/

different scans and between horizontal and vertical polarizations, the calibration uncertainties are estimated to be lower than 10 per cent at 3 mm and 20 per cent in the higher frequency bands. After subtraction of the continuum emission via first-order polynomial fitting, a final spectrum was obtained for each source and frequency band after stitching the spectra from each scan and frequency setting. The intensity was converted from antenna temperature (T_{ant}^*) to main beam temperature (T_{mb}) using the beam efficiencies provided at the IRAM web site.² In order to improve the signal-to-noise ratio (S/N), the 2- and 1-mm ASAI data were smoothed to 0.5 km s⁻¹, except in the case of L1544, for which we kept the original spectral resolution.

For I16293, we used the TIMASSS spectral data obtained with the IRAM 30-m telescope at 1, 2, and 3 mm. A detailed description of the observations and an overview of the data set are reported in Caux et al. (2011b).

4 RESULTS

4.1 Line spectra

We searched for formamide (NH2CHO) and isocyanic acid (HNCO) in our data set using the CASSIS software³ (Caux et al. 2011a) and the Cologne Database for Molecular Spectroscopy (CDMS;⁴ Müller et al. 2001, 2005) to identify the lines. For NH₂CHO, we detected transitions with upper level energies, Eup, below 150 K, and spontaneous emission coefficients, A_{ij} , above 10^{-5} s⁻¹ and 5 × 10^{-5} s⁻¹, respectively, for the 2/3-mm and the 1-mm data. For HNCO, we detected transitions with $E_{\rm u} < 150$ K and $A_{ii} > 10^{-5}$ s⁻¹. Tables B1 and B2 list all the NH₂CHO and HNCO transitions fulfilling these criteria in the observed millimetre bands, as well as the 3σ detections for each source. The sources where no NH2CHO lines were detected (see below) are not included in Table B1. For some sources with not many clear formamide detections (e.g. IRAS4A, Cep E), we included a few additional lines with peak intensities between 2σ and 3σ , as indicated in the tables. We then fitted the lines with a Gaussian function, and excluded from further analysis those falling well below or above the systemic velocity, and/or displaying too narrow or too broad linewidths with respect to the typical values encountered for each source.

Table 2 lists, for each source, the number of NH₂CHO and HNCO lines detected and used in our analysis (Section 4.2). While HNCO is easily detected in all the sources, NH₂CHO remains undetected in five objects: L1544, TMC-1, B1, L1527, and L1157mm. Moreover, in those sources where it is detected, the lines are typically weak (S/N~ 3–5). OMC-2 FIR 4 has the highest number of detected formamide lines, which are also the most intense. The results from the Gaussian fitting to the detected lines are presented in Tables B3–B12. A sample of lines for all the ASAI sources are shown in Figs C1–C3.

4.2 Derivation of physical properties

4.2.1 Rotational diagram analysis

In order to determine the excitation conditions – i.e. excitation temperature, column density and, eventually, abundance with respect to H_2 – of NH₂CHO and HNCO for each source in a uniform way, we employed the CASSIS software to build rotational diagrams.

³ cassis has been developed by IRAP-UPS/CNRS (http://cassis.irap.omp.eu)
⁴ http://www.astro.uni-koeln.de/cdms/

Table 2.	Number of NH2CHO and HNCO detected
lines.	

	NH	I ₂ CHO	HNCO			
Source	No.	$E_{\rm u}$ (K)	No.	$E_{\rm u}$ (K)		
L1544 ^a	0	_	2	10–16		
TMC1	0	-	3	10-16		
B1	0	-	4	10-30		
L1527	0	-	4	10-30		
L1157-mm	0	-	4	10-30		
IRAS 4A	7	15-70	10	10-130		
SVS13A	13	15-130	19	10-130		
OMC-2 FIR 4	21	10-130	9	10-100		
Cep E	5	10-22	5	10-85		
I16293	12	10-160	16	10–95		

Note. ^aOnly 3-mm data available.

This approach assumes (i) that the lines are optically thin, and (ii) Local Thermodynamic Equilibrium (LTE), meaning that a single Boltzmann temperature, known as *rotational temperature*, describes the relative distribution of the population of all the energy levels for a given molecule. Under these assumptions, the upper-level column density

$$N_{\rm u} = \frac{8\pi k v^2}{hc^3 A_{\rm ul}} \frac{1}{\eta_{\rm bf}} \int T_{\rm mb} \mathrm{d}V \tag{1}$$

and the rotational temperature, $T_{\rm rot}$, are related as follows:

$$\ln \frac{N_{\rm u}}{g_{\rm u}} = \ln N_{\rm tot} - \ln Q(T_{\rm rot}) - \frac{E_{\rm u}}{kT_{\rm rot}}$$
(2)

where k, v, h, and c are, respectively, Boltzmann's constant, the frequency of the transition, Planck's constant, and the speed of light; g_u is the degeneracy of the upper level, and N_{tot} is the total column density of the molecule. The second fraction in equation (1) is the inverse of the beam-filling factor. We estimated it assuming sources with a Gaussian intensity distribution:

$$\eta_{\rm bf} = \frac{\theta_{\rm s}^2}{\theta_{\rm s}^2 + \theta_{\rm b}^2} \tag{3}$$

with θ_s and θ_b being, respectively, the source and telescope beam sizes. We adopted the source sizes indicated in Table 3. In those sources where a hot (T > 100 K) inner region is believed to exist, we considered two possible solutions: (i) the emission originates from a compact size representing this inner region or *hot corino*, which typically shows enhanced abundances of COMs; and (ii) the emission homogeneously arises from the entire extended molecular envelope of the protostar. We determined the sizes of the compact hot corino regions either from published interferometric maps (SVS13A) or from the gas density structure, n(r), reported in the literature (I16293, IRAS 4A, OMC-2, Cep E), as indicated in Table 3. In the latter case, we assumed a size equal to the diameter within which the dust temperature is above 100 K.

Some sources, such as IRAS 4A, OMC-2, and Cep E, show extended velocity wings in a few of their lines. In order to separate their contribution to the line emission, we determined their line flux, $\int T_{mb}dV$, by fitting a Gaussian function to the affected lines after masking their high-velocity wings. In sources with two to four well-aligned data points in the rotational diagrams, we took into account the relatively large error bars by fitting two additional 'extreme' lines passing through the tips of the error bars of the lowest and largest energy points. An example is shown for B1 in Fig. 1, where the two extreme solutions are depicted in blue, while

² http://www.iram.es/IRAMES/mainWiki/Iram30mEfficiencies

Table 3. Results from the rotational diagram analysis of NH₂CHO and HNCO: Adopted size and H₂ column densities (N_{H2}), derived rotational temperatures, T_{rot} , derived HNCO and HN₂CHO column densities (N_{HNCO} , N_{NH_2CHO}), resulting abundances with respect to H₂(X_{HNCO} , X_{NH_2CHO}), and ratio of HNCO to NH₂CHO column densities (R).

Source	Size ^a (arcsec)	$N_{\rm H2}^b$ (10 ²² cm ⁻²)	T _{rot(HNCO)} (K)	$\frac{N_{\rm HNCO}}{(10^{12} {\rm cm}^{-2})}$	$\begin{array}{c} X_{\rm HNCO} \\ (10^{-11}) \end{array}$	$T^c_{ m rot(NH_2CHO)}$ (K)	$N_{\rm NH_2CHO}^c$ (10 ¹² cm ⁻²)	$\begin{array}{c} X_{\rm NH_2CHO} \\ (10^{-11}) \end{array}$	R
One-component fit									
L1544	BF	9.4 ± 1.6^{1}	7 ± 3	5 ± 3	5 ± 3	7	< 0.036	< 0.046	>130
$TMC1^d$	BF	1.0 ± 0.1^{2}	4 ± 1	8 ± 5	80 ± 50	4	< 0.47	< 5.2	>17
B1	BF	7.9 ± 0.3^{3}	10 ± 2	8.4 ± 1.6	11 ± 2	10	< 0.087	< 0.11	>97
L1527	BF	4.1^{4}	7.5 ± 1.4	2.5 ± 1.5	6 ± 4	7.5	< 0.062	< 0.15	>40
L1157-mm	30	120^{5}	8 ± 1	4 ± 1	0.35 ± 0.03	8	<1	< 0.008	>40
SVS13A (ext) ^e	20	106	58 ± 6	11 ± 2	11 ± 2	64 ± 6	3.0 ± 0.4	3.0 ± 0.4	4 ± 1
$(\text{com})^e$	1	10007	36 ± 3	1500 ± 300	15 ± 3	40 ± 4	320 ± 60	3.2 ± 0.6	5 ± 1
OMC-2 (ext)	25	19 ⁸	25 ± 3	16 ± 3	1.9 ± 0.4	58 ± 4	3.1 ± 0.2	0.36 ± 0.02	5 ± 1
(com)	2	4.6^{8}	19 ± 1	900 ± 100	910 ± 80	32 ± 2	110 ± 10	110 ± 10	8 ± 1
Cep E (ext)	40	4.89	30 ± 5	6.2 ± 0.3	13 ± 1	9 ± 2	0.2 ± 0.1	0.4 ± 0.2	30 ± 13
(com)	0.5	230 ⁹	17 ± 1	6000 ± 1000	130 ± 15	6 ± 1	500 ± 300	11 ± 5	12 ± 6
Two-component fit									
IRAS 4A (C1)	30	2.9^{10}	11 ± 3	10 ± 1	34 ± 2	19 ± 15	0.6 ± 0.5	1.9 ± 0.2	18 ± 2
IRAS 4A (C2)	0.5	250^{10}	43 ± 8	2000 ± 1000	80 ± 40	30 ± 5	500 ± 100	20 ± 5	4 ± 2
I16293 (C1)	30	2.9^{11}	14 ± 5	20 ± 2	69 ± 7	5 ± 1	1.7 ± 0.6	6 ± 2	12 ± 4
I16293 (C2)	1.2	5311	47 ± 4	4400 ± 700	830 ± 130	83 ± 33	590 ± 190	110 ± 40	8 ± 3

Notes. ^{*a*}BF: beam-filling assumed. For the other sources, the size has been adopted as follows: L1157mm and IRAS 4A (extended) from Jørgensen, Schöier & van Dishoeck (2002); SVS13A (extended) from Lefloch et al. (1998); SVS13A (compact) from Looney, Mundy & Welch (2000); OMC-2 FIR 4 (extended) from Furlan et al. (2014); OMC-2 FIR 4 (compact) from Crimier et al. (2009); Cep E from Chini et al. (2001); IRAS 4A (compact) from Maret et al. (2002); IRAS 16293 from Jaber et al. (2014).

^bReferences: ¹Crapsi et al. (2005), ²Maezawa et al. (1999), ³Daniel et al. (2013), ⁴Parise, Bergman & Menten (2014), ⁵Jørgensen et al. (2002) ⁶Lefloch et al. (1998), ⁷Looney et al. (2000), ⁸Crimier et al. (2009), ⁹Crimier et al. (2010a), ¹⁰Maret et al. (2002), ¹¹Crimier et al. (2010b).

^cFor the non-detections of NH₂CHO, we have computed a 3σ upper limit to its column density adopting the same T_{rot} derived for HNCO (see text).

 $^d\mathrm{Data}$ for $\mathrm{NH_2CHO}$ upper limit derived from 3-mm data by N. Marcelino.

 $^{e}N(\text{HNCO})$ is probably a lower limit due to contamination from the OFF position.

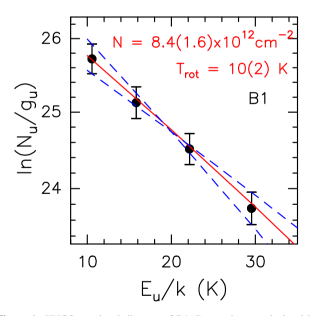


Figure 1. HNCO rotational diagram of B1. Data points are depicted in black. The red lines correspond to the best fit to the data points. The extreme solutions taking into account the error bars are displayed in dashed blue.

the best solution is marked in red. The remaining rotational diagrams and the best fit to their data points using equation (2) are shown in Figs C4 and C5, where the error bars take into account calibration errors as well as the rms value around each line.

We compared our rotational diagram results with those reported in Marcelino et al. (2009) for the four sources common to both studies: L1544, TMC-1, B1, and L1527. The column densities of HNCO are in perfect agreement within the uncertainties, while the rotational temperatures agree within 1 K.

For homogeneity with the methodology used for NH₂CHO, we estimated the properties of HNCO in the LTE approximation. In addition, by adopting the same source sizes for HNCO and NH₂CHO, we assumed that the emission from both molecules originates in the same region(s). The similar average linewidths between the two species suggest this is a reasonable assumption. Table 3 and Figs C4 and C5 present the results of the rotational diagram analysis. For most of the sources, a single component fits well both the NH₂CHO and HNCO points and therefore LTE seems to reproduce well the observations. This can also be seen in Figs C1 to C3, where the observed spectra (in black) and the best-fitting models (in red) match fairly well. However, for SVS13A, Cep E, and OMC-2, the compact solutions correspond to HNCO lines that are moderately optically thick ($\tau \sim 1$ –10). The most extreme case is Cep E, for which also the NH₂CHO lines are optically thick. This is in contradiction with the underlying assumption of optically thin lines in the rotational diagram method. We find, however, that this caveat can be easily overcome by adopting a slightly larger source size, of 3, 2, and 2 arcsec, respectively, for SVS13A, Cep E, and OMC-2. Doing this, the resulting column densities are reduced by a factor of 2 (OMC-2) to 15 (Cep E), τ becomes much smaller than 1, and the lines can be well fitted by the solutions. Consequently, the uncertainties in the compact-solution column densities in these three sources are larger than reported in Table 3, but they are taken into account in the discussion (Section 5: see Figs 2 and 3).

There are two objects where a single component does not appear to explain the emission of all the lines: IRAS 4A and I16293, two

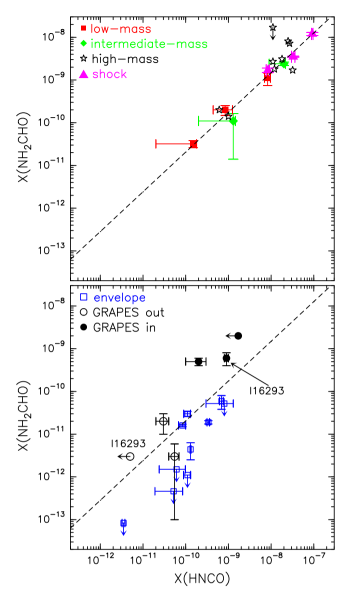


Figure 2. Plot of NH_2CHO versus HNCO abundances with respect to H_2 . Top: data points included in the power-law fit (dashed line; see text). Red squares and green diamonds denote the compact or inner RD solutions of low- and intermediate-mass sources in this study, respectively. Magenta triangles and black stars correspond, respectively, to outflow shock regions (from Mendoza et al. 2014) and high-mass sources (from Bisschop et al. 2007 and Nummelin et al. 2000). Bottom: data points not included in the power-law fit (see text). Blue open squares represent the extended or outer RD solutions, while black open and filled circles denote the GRAPES LTE values for the outer and inner components, respectively.

well-known hot corino sources. Indeed, their rotational diagrams suggest either the contribution of two components, or non-LTE effects, coming into play. Considering the former, Table 3 presents the results of a two-component solution to the rotational diagrams of these two objects, where C1 is assumed to represent the cold extended envelope of the protostar, and C2 the small inner hot corino. While this two-component solution reproduces well the observations, non-LTE effects cannot be ruled out.

As for the five objects where formamide was not detected, we determined a 3σ upper limit to its column density under the assumption of LTE and adopting the corresponding value of $T_{\rm rot}$ derived for HNCO. To this end, we used the spectral data around the NH₂CHO

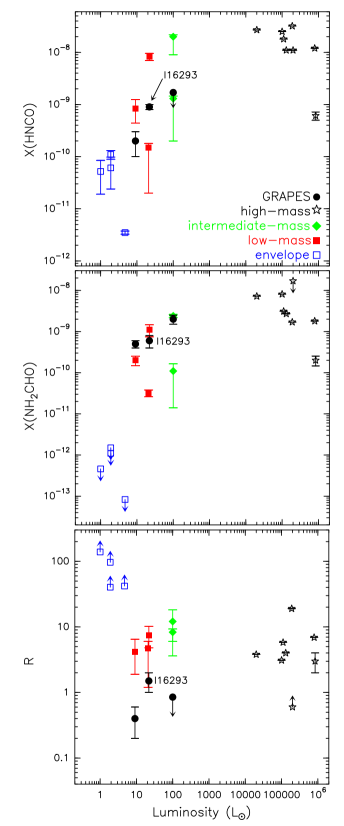


Figure 3. Abundance of HNCO (top), NH_2CHO (middle) and their ratio (bottom) against bolometric luminosity. Symbols are as in Fig. 2.

 $4_{0,4}$ - $3_{0,3}$ transition at 84.542 GHz, expected to be the most intense at the cold temperatures implied by the HNCO results. The upper limits thus derived are shown in Table 3.

Once the column densities of HNCO and NH₂CHO were obtained, we derived their respective abundances with respect to molecular hydrogen (H₂) using the H₂ column densities, N_{H2} , listed in Table 3, which correspond to the indicated source sizes. The uncertainty on N_{H2} is included for those sources where this was provided in the corresponding bibliographic reference. The resulting abundances span more than two orders of magnitude and are shown in Table 3, together with their ratio, $R = X(\text{HNCO})/X(\text{NH}_2\text{CHO})$.

4.2.2 Radiative transfer analysis taking into account the source structure

The source structures of I16293, IRAS 4A, and OMC-2 are reported in the literature (Maret et al. 2002; Crimier et al. 2010b). Therefore, for these objects, a more sophisticated radiative transfer analysis is possible that takes into account the temperature and gas density as a function of distance to the central protostar. Cep E also has a known structure (Crimier et al. 2010a), but having only five line detections, both in HNCO and in NH₂CHO, we do not consider it here. We have analysed the I16293, IRAS 4A, and OMC-2 lines by means of the code GRAPES (GRenoble Analysis of Protostellar Envelope Spectra), whose details are described in Ceccarelli et al. (2003) and Jaber et al. (2014).

Briefly, GRAPES computes the Spectral Line Energy Distribution (SLED) of a free-infalling spherical envelope with given gas and dust density and temperature profiles, and for a given mass of the central object. The dust-to-gas ratio is assumed to be the standard one, 0.01 in mass, and the grains have an average diameter of $0.1 \,\mu m$. The species abundance is assumed to follow a step-function, with a jump at the dust temperature T_{jump} , which simulates the thermal desorption of species from icy mantles (e.g. Ceccarelli et al. 2000). The abundance X_i in the warm ($T \ge T_{jump}$) envelope is constant. In the outer envelope, we assumed that the abundance follows a power law as a function of the radius, $X_0 r^a$, with an index equal to 0, -1 and -2, as in Jaber et al. (2014). X_i and X_o are considered parameters of the model. Since, to our knowledge, the binding energy of NH2CHO is not available in the literature, we treat T_{jump} as a parameter too. However, if the molecules are trapped in water ice, the binding energy of H2O will largely determine the dust temperature at which NH₂CHO is injected into the gas-phase.

The radiative transfer is solved with the escape probability formalism and the escape probability is computed integrating each line opacity over the 4π solid angle. We ran models assuming LTE populations for formamide and, for comparison with Section 4.2.1, HNCO, and models taking into account non-LTE effects for HNCO. In the latter case, we used the collisional coefficients by Green (1986), retrieved from the LAMDA data base (Schöier et al. 2005).

For each molecule and source, we ran a large grid of models varying the four parameters mentioned above: X_i, X_o, T_{jump} , and a. In total, we ran about 20 000 models per source. The computed SLED of each model was then compared with the observed SLED to find the solution with the best fit. The results of this analysis are reported in Table 4, where we give the best-fitting values and the range of X_i, X_o, T_{jump} with $\chi^2 \leq 1$. We note that there is no appreciable difference in the best χ^2 when using a different value of a, so we took the simplest solution: a = 0. In this respect, the situation is similar to what Jaber et al. (2014) found in their study of IRAS16293.

Table 4. Results of grapes analysis for NH_2CHO and HNCO considering the source structure of IRAS 4A, 116293 and OMC-2.*

	IRAS 4A	I16293	OMC-2
HNCO LTE			
$X_{\rm o} (10^{-11})$	3 ± 1	0.1 ± 0.1	5.5 ± 1.5
$X_{\rm i} (10^{-11})$	20 ± 10	90 ± 10	<170
T_{jump} (K)	100	40	80
T_{jump} range (K)	60-120	30-50	≥30
χ^2	1.2	2.0	1.0
HNCO non-LTE			
$X_{\rm o} (10^{-11})$	3 ± 1	0.5 ± 0.4	4 ± 1
X_{i} (10 ⁻¹¹)	30 ± 20	600 ± 300	<20
T_{jump} (K)	100	90	80
T_{jump} range (K)	≥ 50	≥ 60	≥30
χ^2	1.0	1.5	0.7
NH ₂ CHO			
$X_0 (10^{-11})$	2 ± 1	0.3 ± 0.2	0.3 ± 0.3
$X_{\rm i} (10^{-11})$	50 ± 10	60 ± 20	200 ± 50
T _{jump} (K)	100	90	80
T_{jump} range (K)	≥ 100	≥ 50	60-100
χ^2	2.0	0.7	1.3
$R = X(HNCO)/X(NH_2CHO)$			
R_{0} (LTE)	1.5 ± 0.9	<1.7	18 ± 18
R _i (LTE)	0.4 ± 0.2	1.5 ± 0.5	< 0.85
R_{0} (non-LTE)	1.5 ± 0.9	1.7 ± 1.7	13 ± 13
$R_{\rm i}$ (non-LTE)	0.6 ± 0.4	10 ± 6	< 0.1
• • • •			

Note. *Abundances with respect to H₂ are times 10^{-11} . X_0 and X_i are the outer and inner abundances, respectively.

A comparison between the results obtained for HNCO with the LTE and non-LTE level populations shows that the LTE approximation is quite good in the case of IRAS 4A and OMC-2, but not for I16293. The reason for that is probably a lower density envelope of I16293 compared to the other two sources. Therefore, the LTE results are likely reliable also for the formamide in IRAS 4A and OMC-2, while in I16293 these have to be taken with some more caution.

A second result of the GRAPES analysis is that both HNCO and formamide have a jump in their abundances at roughly the same dust temperature, 80–100 K. This is an important result reflecting the two molecules have similar behaviours with changes in temperature. It suggests they trace the same regions within the analysed protostars.

In order to evaluate whether the rotational diagram (hereafter RD) and GRAPES analyses are in agreement, we compare their respective abundance values, which roughly agree within an order of magnitude, in the Appendix A. We note here that, while the GRAPES analysis is likely more accurate, we are not able to apply it to the other sources of this study due to the lack of known source structure and/or lack of a sufficient amount of molecular lines. The absence of interferometric imaging of the HNCO and NH₂CHO emission also hinders the study of the inner structure of the protostellar emission. Therefore, we base the discussion below largely on the RD results, with a note of caution that those values may not strictly represent the physical properties of the sources.

5 DISCUSSION

5.1 Formation routes of NH₂CHO

The formation mechanism(s) of interstellar formamide, as that of other COMs, is still far from being established. Several routes

have been proposed so far which include both gas-phase and grain-surface processes. Concerning the former, Quan & Herbst (2007) suggested NH_2CHO forms via the radiative association reaction

$$NH_4^+ + H_2CO \rightarrow NH_4CH_2O^+ + h\nu$$
(4)

followed by dissociative recombination. Halfen, Ilyushin & Ziurys (2011) proposed the following ion–molecule reaction and subsequent electron recombination:

$$NH_4^+ + H_2CO \rightarrow NH_3CHO^+ + H_2$$
(5)

$$NH_3CHO^+ + e^- \rightarrow NH_2CHO + H.$$
(6)

These reactions all have unknown rates. Thus, further experimental work will be needed in order to evaluate their effectiveness in producing formamide.

Neutral-neutral reactions have also been discussed as possible gas-phase routes leading to NH_2CHO . In particular, Garrod, Weaver & Herbst (2008) proposed the radical-neutral reaction

$$H_2CO + NH_2 \rightarrow NH_2CHO + H.$$
(7)

However, as recently mentioned by Redondo et al. (2014), it presents a net activation barrier of > 1000 K that makes it inviable in interstellar conditions. Other neutral–neutral reactions evaluated by these authors also revealed to have large activation barriers, thus ruling them out as dominant or efficient mechanisms to produce NH₂CHO.

Formamide may also be formed on the icy mantles of dust grains. Jones et al. (2011) conducted some experimental work in which they irradiate a mixture of ammonia (NH₃) and carbon monoxide (CO) ices with high-energy (keV) electrons, resulting in NH₂CHO as one of the final products. The authors discuss several possible reactions and conclude that the most plausible route towards formamide begins with the cleavage of the nitrogen–hydrogen bond of ammonia, forming the NH₂ radical and atomic H. The latter, containing excess kinetic energy, can then add to CO, overcoming the entrance barrier, to produce the formyl radical (HCO). Finally, HCO can combine with NH₂ to yield NH₂CHO.

A different grain-mantle mechanism was proposed by Garrod et al. (2008), who considered hydrogenation (i.e. addition of H atoms) of OCN in their chemical models. However, this route resulted in an overabundance of NH2CHO and an underabundance of HNCO, since the latter was efficiently hydrogenated to yield formamide, the final product. Raunier et al. (2004) performed experimental Vacuum Ultra Violet (VUV) irradiation of solid HNCO at 10 K, which led to NH₂CHO among the final products. They proposed that photodissociation of HNCO yields free H atoms that subsequently hydrogenate other HNCO molecules in the solid to finally give NH₂CHO. The limitation of this experiment is that it was carried out with pure solid HNCO. Jones et al. (2011) mentioned that, in the presence of NH₃, quite abundant in grain mantles, HNCO will preferentially react with it, resulting in $NH_4^+ + OCN^-$. Despite these caveats, hydrogenation of HNCO on grain mantles was recently found to be a most likely solution in the case of the outflow shock regions L1157-B1 and B2 (Mendoza et al. 2014). More experiments and calculations are needed in order to assess the efficiency of this formation route.

5.2 Correlation between HNCO and NH₂CHO

From the previous section, it is clear that, until more gas-phase and surface reaction rates involving the mentioned species are measured,

it will be difficult to establish the exact synthesis mechanisms of formamide in space.

In this section, we assess, from an observational point of view, whether hydrogenation of HNCO leading to NH₂CHO on the icy mantles of dust grains could be a dominant formation route. To this aim, we plot in Fig. 2 the NH2CHO versus HNCO abundances of all our sources (Table 3), as well as the shock regions analysed by Mendoza et al. (2014), and the high-mass SFRs reported in Bisschop et al. (2007) and Nummelin et al. (2000), for comparison. The latter were obtained by the cited authors via the RD method assuming the emission comes from the inner hot core regions. Thus, for homogeneity, we split the plot into two panels, the upper one showing only the compact/inner solutions of the RD analysis, classified by masses. The best power-law fit to these points is marked with a dashed line, and is given by the equation $X(NH_2CHO) = 0.04X(HNCO)^{0.93}$. with a Pearson coefficient of 0.96, indicating a tight correlation. The fact that this correlation is almost linear and holds for more than three orders of magnitude in abundance suggests that HNCO and NH₂CHO are chemically related. This result confirms, on a more statistical basis, what was recently found by Mendoza et al. (2014).

However, this correlation does not hold for the objects without formamide detections, which are plotted in the lower panel of Fig. 2 together with the extended envelope solutions of the RD analysis. Here, it is clearly seen that all the upper limits lie well below the bestfitting line, indicating a significantly larger amount of gas-phase HNCO relative to NH₂CHO in comparison to the other sources. These objects are the coldest in our sample, representing either prestellar cores or protostars with no detectable hot corino within them. The rotational temperatures inferred from the HNCO RD analysis are also among the lowest in our sample. In this same plot, the points representing formamide detections (extended envelope component) also show a tendency towards lower relative values of $X(NH_2CHO)$, although not as pronounced.

Thus, it appears that regions with colder temperatures are more deficient in NH₂CHO than protostars with hot inner regions, indicating that higher temperatures are needed for NH2CHO to become relatively abundant in the gas-phase. This might be explained by (i) NH₂CHO forming in the gas-phase at temperatures above \sim 100 K, and/or (ii) it forming predominantly on the icy mantles of dust grains at low temperatures, and subsequently sublimating into the gas-phase when the temperature in the inner regions rises sufficiently. As for the former possibility, Mendoza et al. (2014) quantitatively argued that reaction 7 does not suffice to explain the amount of gas-phase formamide in the shock regions of L1157 protostellar outflow. In addition, the high activation barrier the reaction needs to overcome makes this an unviable route. Other purely gas-phase formation routes still need more investigation in terms of reaction rates and activation barriers, as discussed in Section 5.1. Mendoza et al. (2014) favoured a grain formation mechanism followed by mantle-grain evaporation/sputtering on the basis of the comparable abundance enhancements of HNCO, NH₂CHO, and CH₃OH in the gas-phase between the two protostellar shocks studied by the authors. Therefore, grain formation of NH₂CHO appears to be the most likely possibility.

On the other hand, while grain formation of HNCO is likely to occur in the cold phases of star formation (Hasegawa & Herbst 1993), gas-phase reactions leading to HNCO at such cold temperatures can also take place efficiently (see e.g. Marcelino et al. 2009 and references therein), overcoming strong depletion. This would explain its relatively high gas-phase abundance already in the very early – and cold – phases of star formation, and also the high

values of HNCO to NH₂CHO abundance ratios we find in the coldest sources of our sample.

In Fig. 3, we plot the HNCO abundance, the NH₂CHO abundance, and their ratio, R, as a function of bolometric luminosity for those sources with a reported luminosity estimate (see Table 1). For the objects in our study with formamide detection, we only plot the points corresponding to the inner or compact component (red circles), since these regions are expected to be the dominant contributors to the overall luminosity. The HNCO and NH₂CHO abundance panels both show the high-mass sources lying on top of the plot, while the points representing our sample sources are more scattered, with the coldest objects (in blue) showing the lowest abundances. This trend is much more pronounced in the case of NH₂CHO, for which hot corino regions (red points) display higher NH₂CHO abundances than the colder objects by more than an order of magnitude. More interesting is the plot of R, which illustrates how this quantity remains roughly constant along 6 orders of magnitude in luminosity for the NH2CHO-emitting sources, with values ranging from 3 to 10 approximately. This reflects the almost-linearity of the correlation between the abundance of the two species. On the other hand, this value rises considerably for the lower luminosity sources, re-enforcing our interpretation that formamide mostly forms on grains at cold temperatures, while HNCO may form both on grains and in the gas.

The strikingly tight and almost linear correlation between the abundance of the two molecules once NH_2CHO becomes detectable suggests one of the two following possibilities: (i) HNCO and NH_2CHO are both formed from the same parent species on dust grain mantles, or (ii) one forms from the other. Among the grain formation routes that have been proposed so far, hydrogenation of HNCO leading to NH_2CHO is the only mechanism that would explain our observational results. While this route is found to have some caveats (see Section 5.1), it is also true that more experimental work is needed to better assess its efficiency.

If the abundance of gaseous NH_2CHO truly depends on temperature, we should find a difference in *R* between the hot corino and the cold envelope regions of IRAS 4A and I16293. Looking at Table 3, this is indeed the case. As for OMC-2, Cep E, and SVS13A, only one component was necessary to describe their rotational temperatures and column densities. Therefore, we cannot compare the extended and compact values as in the case of a two-component solution. We can nevertheless guess that, excluding the case of Cep E, for which only low-energy formamide lines were detected, the compact solution is likely the best, given the low values of *R* and the relatively high rotational temperatures derived. This would imply most of the emission arises in the inner hot corino regions. For Cep E, more molecular observations at higher frequencies are needed to confirm this.

Figs 2 and 3 also include the results from the LTE GRAPES analysis. I16293 (labelled in the plots) is included for completeness despite the fact that the GRAPES analysis suggests non-LTE effects should be taken into account for this object. While these points introduce more scatter in the plots, it can be clearly seen that the inner components of the sources analysed with GRAPES have a lower HNCO abundance relative to NH₂CHO, compared to what is found via the RD analysis. This yields lower *R* values, indicating a considerable amount of formamide with respect to HNCO in these regions and suggesting, as mentioned in Section A, that the two-component approximation in the RD analysis is oversimplistic: while we assumed that only the higher-energy formamide lines arose from the compact inner region, it is likely that a significant amount of emission from the

low-energy lines also originates here and not exclusively in the outer envelope.

The trend showing higher R in the outer envelope than in the inner regions holds for both IRAS 4A and OMC-2, which further supports the fact that NH₂CHO requires higher temperatures than HNCO to be detectable in the gas-phase. This kind of analysis, taking into account the source structure, is needed in a larger sample of objects in order to draw conclusions about both the chemistry and the validity of our RD analysis on a more statistical basis. Interferometric mapping would also greatly help disentangling source multiplicity and verifying whether the emission of HNCO and NH₂CHO trace the same regions, as has been assumed in this work.

6 CONCLUSIONS

As part of the IRAM Large programme ASAI, we searched for millimetre spectral lines from formamide (NH₂CHO), a presumably crucial precursor of pre-biotic material, and isocyanic acid (HNCO), in 10 low- and intermediate-mass SFRs with different properties. The data set, obtained with the IRAM 30-m telescope, consists mainly of unbiased broad-band spectral surveys at 1, 2, and 3 mm. Our aim was to investigate the chemical connection between these two molecular species and gain some observational insights into the formation mechanisms of formamide in interstellar conditions. The present work represents the first systematic study within ASAI and statistically completes the low-mass end of similar studies performed towards high-mass SFRs. Our main findings are summarized as follows.

(1) The high sensitivity and large frequency range of the spectral surveys allowed us to evaluate the detectability of numerous NH_2CHO and HNCO transitions. We detect formamide in five out of the 10 objects under study (IRAS 4A, IRAS 16293, SVS13A, Cep E, and OMC-2), and HNCO in all of them. Since formamide had already been detected in IRAS 16293 – also investigated here for completeness – this study raises the number of known low-and intermediate-mass formamide-emitting protostars to five, thus significantly improving the statistics.

(2) We derived HNCO and NH₂CHO column densities via the rotational diagram method for all the sources. As a result, we found NH₂CHO abundances with respect to H₂ in the range 10^{-11} – 10^{-9} , and HNCO abundances between 10^{-12} and 10^{-8} . For those objects without formamide detection, we provided an upper limit to its column density and abundance.

(3) For three targets (IRAS 4A, IRAS 16293, and OMC-2), the source density and temperature structures are known and published, and we were thus able to take them into account through a more sophisticated analysis using the code GRAPES. This method fits an abundance profile that consists of a step function, with the separation between the two values roughly corresponding to the hot corino size. A comparison between the two radiative transfer analyses employed reveals overall agreement within an order of magnitude. The GRAPES analysis also indicates that one of the studied objects, IRAS 16293, requires a non-LTE radiative transfer analysis, which at the moment is not possible due to the lack of collisional coefficients for NH₂CHO. LTE appears to describe correctly the other two sources analysed with GRAPES, and is assumed to be a good approximation for all the other sources in our sample.

(4) For the sources where formamide was detected, i.e. hot corino sources, we found an almost linear correlation between HNCO and NH₂CHO abundances that holds for several orders of magnitude. This suggests that the two molecules may be chemically associated.

On the other hand, those sources with no formamide detection do not follow this correlation, but instead show much larger amounts of HNCO relative to NH_2CHO . These objects are the coldest in this study, and unlike the rest of our sample, they contain no known hot corinos.

(5) Our findings and the NH₂CHO formation routes proposed so far in the literature suggest that, unlike HNCO, NH₂CHO does not form efficiently in the gas-phase at cold temperatures and may be formed on the mantles of dust grains, where it remains frozen at cold temperatures. As soon as the temperature rises sufficiently to sublimate the icy grain mantles, formamide is incorporated into the gas and becomes detectable. The tight and almost linear correlation with HNCO suggests a possible formation route of NH₂CHO via hydrogenation of HNCO, although other possibilities should not be ruled out. In particular, two potentially viable gas-phase pathways leading to formamide involve formaldehyde (H₂CO). It is therefore worth exploring the connection between H₂CO and NH₂CHO, which will be the subject of a forthcoming paper.

(6) In order to evaluate the validity of our conclusions, several aspects need to be explored more thoroughly. From an observational point of view, interferometric imaging is necessary to assess the relative spatial distribution of HNCO and NH₂CHO, and retrieve more accurate abundance ratios, in particular in the hot corino sources. In addition, more detailed and sophisticated radiative transfer analysis requires, on the one hand, knowledge of the source density and temperature profiles and, on the other hand, collisional coefficient calculations for NH₂CHO, currently unavailable. Finally, more chemical experiments are needed to estimate the efficiency of the hydrogenation processes leading from isocyanic acid to formamide on interstellar dust grains, as well as the viability of purely gas-phase reactions.

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APPENDIX A: COMPARISON BETWEEN GRAPES AND ROTATIONAL DIAGRAM ANALYSES

This section aims to compare the agreement between the rotational diagram and GRAPES methods. As described in Section 4.2.2, the line emission in I16293 does not appear to be well described by LTE, and a more realistic radiative transfer treatment will need to wait until collisional coefficients are available for NH₂CHO. Therefore, we do not consider it here, while it is worth noticing that a rotational diagram analysis is likely too simplistic to analyse the HNCO and NH₂CHO lines in this source.

In OMC-2, the GRAPES analysis tells us that the temperature that separates the inner and outer components is 80 K, in both the LTE and non-LTE approximations. Thus, for consistency in the comparison, we re-computed the inner and outer abundances resulting from the RD analysis using the same inner sizes as in GRAPES, instead of those corresponding to a temperature of the 100 K (see Section 4.2.1). We note that, while the RD analysis allowed for a separation of two components (inner and outer) for IRAS 4A, a single component was sufficient for OMC-2. It should be kept in mind, therefore, that for the latter the comparison is not equivalent, since we are not comparing a two-component solution with another two-component solution as in the case of the other two protostars.

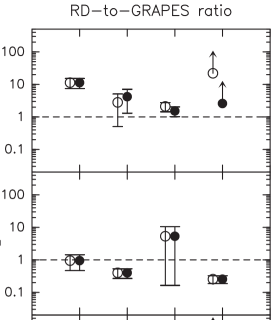
The results of the comparison are listed in Table A1 and illustrated in Fig. A1, where we present the comparison using both the LTE and non-LTE results from GRAPES. It is evident that LTE and non-LTE yield practically the same results for these two sources. It can also be seen that the errors are quite high in some cases, up to 100 per cent, which are caused by the large uncertainties resulting from the GRAPES analysis. Taking these into account, we find the following behaviours:

(i) HNCO abundance: generally, both methods agree within an order of magnitude, but there is a tendency towards higher values in the RD analysis, by a factor of a few.

Table A1. Comparison between ${}_{\mbox{\scriptsize GRAPES}}$ and RD analyses.*

	IRAS 4A	OMC-2
Inner size (arcsec)	1.5	3.1
RD-to-grapes ratio (LTE)		
X _o (HNCO)	11 ± 4	1.5 ± 0.5
X _i (HNCO)	4 ± 3	>2.5
$X_{o}(NH_{2}CHO)$	1 ± 1	5 ± 5
$X_i(NH_2CHO)$	0.4 ± 0.4	0.3 ± 0.3
Ro	12 ± 7	0.3 ± 0.3
Ri	11 ± 8	>10
RD-to-grapes ratio (non-LTE)		
X _o (HNCO)	11 ± 4	2.1 ± 0.7
X _i (HNCO)	3 ± 2	>22
$X_{o}(NH_{2}CHO)$	1 ± 1	5 ± 5
$X_i(NH_2CHO)$	0.4 ± 0.4	0.3 ± 0.3
Ro	12 ± 7	0.4 ± 0.4
Ri	7 ± 6	>85

Note. *Abundances with respect to H_2 are times 10^{-11} .



X(HNCO)

х(NH₂CHO)

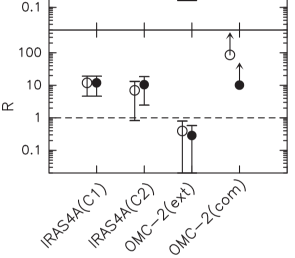


Figure A1. Ratio of RD-to-GRAPES abundances. Top: HNCO abundance. Middle: NH₂CHO abundance. Bottom: HNCO to NH₂CHO abundance ratio. Filled and open circles represent, respectively, the LTE and non-LTE HNCO solution in the GRAPES analysis. The horizontal dashed lines mark equality between RD and GRAPES values.

(ii) NH₂CHO abundance: again, we find agreement within a factor of a few. The compact solution is systematically lower in the RD treatment. This suggests that a non-negligible amount of emission from low-energy molecular lines actually comes from the inner region, and not exclusively from the extended envelope, as assumed in the linear fitting of the RD. Such a finding reflects the necessity of analysis like that performed with GRAPES if we want to properly disentangle the inner and outer components in hot corino or hot core sources.

(iii) HNCO to NH_2CHO abundance ratio, *R*: in this case, the two analysis methods agree within a factor of a few.

APPENDIX B: TABLES

Transition	v (MHz)	<i>E</i> _u (K)	$A_{\rm ul} \ (10^{-5} \ {\rm s}^{-1})$	$\theta_{\rm b}$ (arcsec)	OMC-2	CepE	SVS13A	IRAS4A	I16293	Blends
41,4-31,3	81 693.446	12.8	3.46	30	N	Y*	N	N	N	
40,4-30,3	84 542.330	10.2	4.09	29	Y	Y	Ν	Ν	Y	
42,3-32,2	84 807.795	22.1	3.09	29	Ν	Ν	Ν	Ν	Ν	
43,2-33,1	84 888.994	37.0	1.81	29	Ν	Ν	Ν	Ν	Ν	
43,1-33,0	84 890.987	37.0	1.81	29	Ν	Ν	Ν	Ν	Ν	
$4_{2,2}-3_{2,1}$	85 093.272	22.1	3.13	29	Ν	Y	Ν	Ν	Ν	
$4_{1,3}-3_{1,2}$	87 848.873	13.5	4.30	28	Y	\mathbf{Y}^*	Ν	Ν	Y	
$5_{1,5}-4_{1,4}$	102 064.267	17.7	7.06	24	Y	Y	Ν	В	Y	H ₂ COH ⁺ ?
$5_{0,5}-4_{0,4}$	105 464.219	15.2	8.11	23	Y	Y	Y	Y	Y	2
5 _{2,4} -4 _{2,3}	105 972.599	27.2	6.92	23	Y	N	N	N	N	
5 _{4,2} -4 _{4,1}	106 107.870	63.0	2.98	23	N	N	В	В	N	NH ₂ CHO
5 _{4,1} -4 _{4,0}	106 107.895	63.0	2.98	23	N	N	B	B	N	NH ₂ CHO
5 _{3,3} -4 _{3,2}	106 134.427	42.1	5.29	23	N	N	N	N	N	10120110
5 _{3,2} -4 _{3,1}	106 141.400	42.1	5.29	23	N	N	N	N	N	
$5_{2,3}-4_{2,2}$	106 541.680	27.2	7.03	23	N	N	N	Y	N	
$5_{2,3} + 2,2$ $5_{1,4} - 4_{1,3}$	109 753.503	18.8	8.78	22	N	N	N	N	N	
$5_{1,4} - 4_{1,3}$ $6_{1,5} - 5_{1,4}$	109 755.505 131 617.902	25.1	15.6	19	Y	_	N	N	N	
$7_{1,7}-6_{1,6}$	142 701.325	30.4	20.2	17	Y	_	Y	Y	N	
	142 701.323 146 871.475	28.3	20.2	17	B	_	B	B	B	CU OCU
7 _{0,7} –6 _{0,6}	140 871.473	40.4	22.3	17	B	_	ы N	D N	B	CH ₃ OCH ₃ HCNH ⁺
7 _{2,6} -6 _{2,5}										ICNI '
76,1-66,0	148 555.852	135.7	6.18	17	N	-	N	N	N	
7 _{6,2} -6 _{6,1}	148 555.852	135.7	6.18	17	N	-	N	N	N	
75,3-65,2	148 566.822	103.0	11.4	17	N	-	N	N	N	
75,2-65,1	148 566.823	103.0	11.4	17	N	-	N	N	N	
74,4-64,3	148 598.970	76.2	15.7	17	N	-	N	N	N	
74,3-64,2	148 599.354	76.2	15.7	17	N	-	N	N	N	
73,5-63,4	148 667.301	55.3	19.1	17	Y	-	N	N	N	
73,4-63,3	148 709.018	55.4	19.1	17	Y	-	Y	N	N	
76,2-66,1	149 792.574	40.6	21.9	16	Ν	-	Ν	Ν	\mathbf{Y}^*	
7 _{1,6} –6 _{1,5}	153 432.176	32.5	25.1	16	Y	-	Ν	Ν	Ν	
$13_{2,11} - 13_{1,12}$	155 894.300	105.9	1.26	16	Ν	-	Ν	Ν	Ν	
$12_{2,10} - 12_{1,11}$	155 934.098	92.4	1.23	16	Ν	-	Ν	Ν	Ν	
$11_{2,9} - 11_{1,10}$	157 072.457	79.9	1.22	16	Ν	-	N	Ν	Ν	
$14_{2,12} - 14_{1,13}$	157 115.035	120.5	1.32	16	Ν	-	N	Ν	Ν	
$10_{2,8} - 10_{1,9}$	159 127.569	68.4	1.21	15	-	-	Ν	Ν	Ν	
152,13-151,14	159 739.080	136.2	1.39	15	-	-	Ν	Ν	Ν	
9 _{2,7} -9 _{1,8}	161 899.774	58.1	1.22	15	-	_	Ν	Ν	Ν	
81,8-71,7	162 958.657	38.2	30.5	15	-	-	Ν	Y^*	Y	
82,6-81,7	165 176.756	48.8	1.22	15	-	-	Ν	Ν	Ν	
80,8-70,7	167 320.697	36.4	33.5	15	-	-	Ν	Y^*	Y	
$7_{2,5} - 7_{1,6}$	168 741.408	40.6	1.23	15	-	-	Ν	Ν	Ν	
82,7-72,6	169 299.154	48.5	32.6	15	-	-	Ν	Ν	Ν	
86,2-76,1	169 790.683	143.9	15.3	14	_	_	Ν	Ν	Ν	
86,3-76,2	169 790.683	143.9	15.3	14	_	_	Ν	Ν	Ν	
85,4-75,3	169 810.709	111.1	21.4	14	_	_	Ν	Ν	Ν	
85,3-75,2	169 810.715	111.1	21.4	14	_	_	Ν	Ν	Ν	
84,5-74,4	169 861.469	84.3	26.3	14	_	_	Ν	Ν	Ν	
84,4-74,3	169 862.523	84.3	26.3	14	_	_	Ν	Ν	Ν	
83,6-73,5	169955.835	63.5	30.2	14	_	_	N	N	N	
8 _{3.5} -7 _{3.4}	170039.076	63.5	30.3	14	_	_	N	N	Y	
8 _{2,6} -7 _{2,5}	171620.760	48.8	33.9	14	-	_	N	N	N	
62,4-61,5	172381.012	33.4	1.24	14	_	_	Ν	Ν	Ν	
$10_{1,10} - 9_{1,9}$	203335.761	56.8	60.3	12	Y	_	Ν	Ν	\mathbf{Y}^*	
$10_{0,10} - 9_{0,9}$	207679.189	55.3	64.7	12	Y	_	Ν	\mathbf{Y}^*	Ν	
$10_{2,9} - 9_{2,8}$	211328.960	67.8	65.6	12	Y	_	Y	Y	\mathbf{Y}^b	
105,6-95,5	212323.555	130.5	52.0	12	N	_	N	N	N	
105.6-255										

Table B1. NH₂CHO transitions searched for in this study and 3σ detections.^{*a*}

Table B1. - continued

Transition	v (MHz)	<i>E</i> u (K)	$A_{\rm ul} \ (10^{-5} {\rm s}^{-1})$	$\theta_{\rm b}$ (arcsec)	OMC-2	CepE	SVS13A	IRAS4A	I16293	Blends
104.7-94.6	212428.020	103.7	58.4	12	Ν	_	Ν	Ν	В	NH ₂ CHO
$10_{4.6} - 9_{4.5}$	212433.449	103.7	58.4	12	Ν	_	Ν	Ν	В	NH ₂ CHO
103.8-93.7	212572.837	82.9	63.3	12	Y	_	Y	Ν	Ν	
103.7-93.6	212 832.307	82.9	63.6	12	Ν	_	Ν	Ν	Ν	
102.8-92.7	215 687.009	68.4	69.8	11	Y	_	Ν	Ν	Y	
$10_{1.9} - 9_{1.8}$	218 459.213	60.8	74.7	11	В	_	Ν	Ν	Ν	CH ₃ OH
$11_{1.11} - 10_{1.10}$	223 452.512	67.5	80.5	11	Y	_	Ν	Ν	Ν	
110,11-100,10	227 605.658	66.2	85.5	11	Y	_	Ν	Ν	Ν	
$11_{2,10} - 10_{2,9}$	232 273.646	78.9	88.2	11	Ν	_	Ν	Ν	Y	
115.7-105.6	233 594.501	141.7	73.6	11	Ν	_	В	Ν	Ν	
115,6-105,5	233 594.501	141.7	73.6	11	Ν	_	В	Ν	Ν	
$11_{4,8} - 10_{4,7}$	233 734.724	114.9	80.7	11	Ν	_	Y	Ν	Ν	
114,7-104,6	233 745.613	114.9	80.7	11	Ν	_	Y	Ν	Ν	
113.9-103.8	233 896.577	94.1	86.2	11	Y	_	Y	Ν	Y	
113.8-103.7	234 315.498	94.2	86.7	10	_	_	Ν	Ν	Ν	
$11_{2,9} - 10_{2,8}$	237 896.684	79.9	94.8	10	Y	_	Y	Ν	В	?
$11_{1.10} - 10_{1.9}$	239 951.800	72.3	99.6	10	Y	_	Y	Ν	Ν	
121,12-111,11	243 521.044	79.2	105	10	Ν	_	В	Ν	В	CH ₂ DOH
$12_{0.12} - 11_{0.11}$	247 390.719	78.1	110	10	Ν	_	Ν	Ν	Ν	
$12_{2,11} - 11_{2,10}$	253 165.793	91.1	115	10	Y	_	Ν	Ν	Ν	
$12_{4,9} - 11_{4,8}$	255 058.533	127.2	108	10	Y*	_	Y*	Ν	Ν	
124,8-114,7	255 078.912	127.2	108	10	Ν	_	Ν	Ν	Ν	
$12_{3,10} - 11_{3,9}$	255 225.651	106.4	114	10	Y	_	Y	Ν	Ν	
123,9-113,8	255 871.830	106.4	115	10	Y	_	Ν	Ν	Ν	
$12_{2,10} - 11_{2,9}$	260 189.090	92.4	125	9	В	_	В	Ν	В	H_2C_2O
$12_{1,11} - 11_{1,10}$	261 327.450	84.9	129	9	Ν	_	Y	Ν	Ν	
$13_{1,13} - 12_{1,12}$	263 542.236	91.8	133	9	Y	-	Y	Ν	Ν	

Notes. ^{*a*} Y: Detected above $T_{mb} = 3\sigma$. Y^{*}: Weakly detected (S/N ~ 2–3; see Section 4.1). N: undetected. B: possibly detected but blended. —: not observed.

^bDetected but with an anomalously high flux (maybe blended): removed from analysis.

Table B2. HNCO transitions searched for in this study and 3σ detections.^{*a*}

Transition	v (MHz)	<i>E</i> _u (K)	$A_{\rm ul} (10^{-5} {\rm s}^{-1})$	$\theta_{\rm b}$ (arcsec)	OMC-2	CepE	SVS13A	IRAS4A	I16293	L1157	L1527	B1	L1544	TMC-1
41,4-31,3	87 597.330	53.8	0.80	28	Ν	Ν	Y	\mathbf{Y}^{b}	Ν	Ν	Ν	Ν	Ν	_
40,4-30,3	87 925.237	10.5	0.88	28	Y	Υ	Y	Y	Y	Y	Y	Y	Y	-
41,3-31,2	88 239.020	53.9	0.82	28	\mathbf{Y}^*	Ν	\mathbf{Y}^*	\mathbf{Y}^{b}	Ν	Ν	Ν	Ν	Ν	-
51,5-41,4	109 495.996	59.0	1.7	22	Ν	Ν	Y	Ν	Y	W	Ν	Ν	Ν	Ν
50,5-40,4	109 905.749	15.8	1.8	22	Y	Y	Y	Y	Y	Y	Y	Y	Y	Y
51,4-41,3	110 298.089	59.2	1.7	22	Y^*	Ν	Y	Ν	Y	Ν	Ν	Ν	Ν	Ν
61,6-51,5	131 394.230	65.3	2.9	19	Ν	Ν	Y	Ν	Y	Ν	Ν	Ν	-	Ν
60,6-50,5	131 885.734	22.2	3.1	19	Y	Y	Y	Y	Y	Y	Y	Y	-	Y
61,5-51,4	132 356.701	65.5	3.0	19	Ν	Ν	Ν	Ν	Y	Y	Ν	Ν	_	Ν
71,7-61,6	153 291.935	72.7	4.7	16	Ν	_	Y	Ν	Y	Ν	Ν	Ν	-	Ν
70,7-60,6	153 865.086	29.5	4.9	16	Y	_	Y	Y	Y	Y	Y	Y	_	Y
71,6-61,5	154 414.765	72.9	4.8	16	Ν	_	Y	Ν	Y	Ν	Ν	Ν	-	Ν
101,10-91,9	218 981.009	101.1	14.2	11	Ν	Ν	Y	Ν	Y	Ν	Ν	Ν	-	-
100,10-90,9	219 798.274	58.0	14.7	11	Y	Y	Y	Y	Y	Ν	Ν	Y	-	-
$10_{1,9} - 9_{1,8}$	220 584.751	101.5	14.5	11	Y	Ν	Y	Y	Y	Ν	Ν	Ν	-	-
$11_{1,11} - 10_{1,10}$	240 875.727	112.6	19.0	10	Ν	_	Y	Y	Y	Ν	Ν	Ν	_	-
110,11-100,10	241 774.032	69.6	19.6	10	В	_	В	В	В	Ν	Ν	Ν	-	-
$11_{1,10} - 10_{1,9}$	242 639.704	113.1	19.5	10	Ν	-	Y	Ν	Ν	Ν	Ν	-	_	
121,12-111,11	262 769.477	125.3	24.8	9	Ν	Ν	Y	Y	\mathbf{Y}^{c}	Ν	Ν	Ν	-	-
120,12-110,11	263 748.625	82.3	25.6	9	Y	Y	Y	Y	Y	Ν	Ν	Ν	-	_
$12_{1,11} - 11_{1,10}$	264 693.655	125.9	25.4	9	Ν	Ν	Y	Y	Y	Ν	Ν	Ν	-	-

Notes. ^{*a*}Y: Detected above $T_{\rm mb} = 3\sigma$. Y^{*}: Weakly detected (S/N $\sim 2-3$; see Section 4.1). N: undetected. B: detected but blended. —: not observed.

^bDetected but with an anomalously high flux (maybe blended): removed from analysis.

^cBlended with an unidentified feature: removed from analysis.

Blends: CH₃OH at 241.774 GHz

Table B3. L1544: Gaussian fits to the detected HNCO lines.

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV (\rm mK \ km \ s^{-1})$
$\begin{array}{c} 4_{0,4} - 3_{0,3} \\ 5_{0,5} - 4_{0,4} \end{array}$	87925.237	11.9	459 (2)	7.2 (1)	0.7 (2)	342 (9)
	109905.996	4.8	601 (2)	7.6 (1)	0.7 (1)	448 (5)

Table B4. TMC-1: Gaussian fits to the detected HNCO lines.

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV (\rm mK \ km \ s^{-1})$
50,5-40,4	109 905.996	6.8	157 (8)	5.8 (1)	1.2 (1)	203 (11)
60,6-50,5	131 885.734	6.7	94 (7)	5.8 (1)	0.5 (3)	50 (5)
70,7-60,6	153 865.086	3.2	28 (3)	5.9 (1)	0.5 (4)	15 (2)

Table B5. B1: Gaussian fits to the detected HNCO lines.

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV (\rm mK \ km \ s^{-1})$
40,4-30,3	87 925.237	3.0	530 (12)	6.7 (1)	1.4 (1)	765 (5)
50,5-40,4	109 905.996	22.2	480 (6)	6.6 (1)	1.3 (1)	662 (32)
60,6-50,5	131 885.734	6.3	345 (7)	6.7 (1)	1.4 (1)	521 (8)
70,7-60,6	153 865.086	7.4	224 (6)	6.6 (1)	1.4 (1)	326 (10)

Table B6. L1527: Gaussian fits to the detected HNCO lines.

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV (\rm mK \ km \ s^{-1})$
40,4-30,3	87 925.237	2.3	145 (2)	5.9 (1)	1.3 (1)	198 (3)
$5_{0.5} - 4_{0.4}$	109 905.996	7.9	135 (7)	5.9(1)	1.2 (1)	175 (11)
60,6-50,5	131 885.734	6.7	138 (1)	5.9 (1)	0.8 (1)	115 (8)
70,7-60,6	153 865.086	8.0	63 (1)	5.8 (1)	0.7 (1)	47 (6)

 Table B7.
 L1157mm: Gaussian fits to the detected HNCO lines.

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV (\rm mK \ km \ s^{-1})$
40,4-30,3	87 925.237	3.3	113 (7)	2.6 (1)	1.6 (1)	198 (7)
50,5-40,4	109 905.996	7.8	142 (5)	2.5 (1)	1.2 (1)	177 (13)
60,6-50,5	131 885.734	5.5	81 (3)	2.6 (1)	1.5(1)	126 (11)
70,7-60,6	153 865.086	5.3	71 (1)	2.6 (1)	1.1 (1)	81 (8)

Table B8. IRAS 4A: Gaussian fits to the detected NH₂CHO and HNCO lines.

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV$ (mK km s ⁻¹)
			NH ₂ CHO			
50,5-40,4	105 464.219	3.3	9.8 (1)	6.7 (4)	4.0 (13)	41 (11)
52,3-42,2	106 541.680	2.2	7.7 (17)	7.6 (3)	2.8 (9)	23 (5)
71,7-61,6	142 701.325	5.0	15 (4)	8.1 (3)	2.6 (5)	40 (8)
$8_{1,8} - 7_{1,7}^{w}$	162 958.657	38.2	14 (6)	8.5 (4)	2.9 (7)	44 (9)
$8_{0.8} - 7_{0.7}^{W}$	167 320.697	36.4	13 (4)	7.2 (4)	3.2 (7)	43 (10)
$10_{0,10} - 9_{0,9}^{W}$	207 679.189	55.3	19 (10)	6.6 (5)	3.3 (10)	66 (19)
102,9-92,8	211 328.960	7.1	31 (3)	6.7 (2)	2.4 (5)	82 (15)
			HNCO			
40,4-30,3	87 925.237	3.1	195 (16)	7.2 (1)	2.2(1)	458 (6)
$5_{0,5}-4_{0,4}$	109 905.749	8.0	198 (15)	7.1 (1)	2.3 (1)	495 (16)
60,6-50,5	131 885.734	5.9	203 (12)	7.0(1)	2.5 (1)	545 (11)
70.7-60.6	153 865.086	6.8	168 (11)	7.1 (1)	2.2 (1)	395 (12)
$10_{0,10} - 9_{0,9}$	219 798.274	9.0	88 (8)	6.8 (1)	3.3 (3)	307 (19)
$10_{1,9} - 9_{1,8}$	220 584.751	6.8	37 (6)	6.5 (3)	2.8 (7)	111 (22)
$11_{1,11} - 10_{1,10}$	240 875.727	7.2	23 (2)	6.6 (2)	1.7 (6)	42 (13)
$12_{1,12} - 11_{1,11}$	262 769.477	8.9	44 (4)	6.6 (2)	3.8 (6)	177 (21)
120,12-110,11	263 748.625	12.9	54 (4)	6.5 (8)	3.9 (29)	230 (130)
$12_{1,11} - 11_{1,10}$	264 693.655	9.1	26 (5)	6.3 (4)	3.7 (9)	103 (21)

Note. ^w Transition weakly detected (see Table B1) but included in the analysis for completeness.

Table B9. 116293: Gaussian fits to the detected NH₂CHO and HNCO lines (intensity in T_{ant}^* units).

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm a} {\rm d}V$ (mK km s ⁻¹)
			NH ₂ CHO			
40,4-30,3	84 542.330	5.3	17 (6)	3.2 (12)	6.7 (20)	120 (40)
41,3-31,2	87 848.873	2.6	12 (4)	2.6 (5)	3.8 (11)	50 (13)
51,5-41,4	102 064.267	3.3	16 (5)	2.0 (3)	2.4 (9)	42 (12)
50,5-40,4	105 464.219	5.3	20(7)	5.1 (4)	3.0 (10)	64 (19)
7 _{6,2} -6 _{6,1} ^w	149 792.574	10.0	29 (10)	2.6 (4)	3.0(7)	93 (22)
81,8-71,7	162 958.657	10.1	26 (10)	2.8 (4)	2.5 (13)	69 (27)
80,8-70,7	167 320.697	10.6	55 (11)	2.0 (2)	2.3 (5)	136 (24)
83,5-73,4	170 039.076	15.3	50 (80)	2.7 (17)	2.4 (46)	120 (170)
$10_{1,10} - 9_{1,9}^{W}$	203 335.761	6.8	19 (7)	0.5 (10)	6.7 (27)	135 (43)
102,8-92,7	215 687.009	5.5	30 (10)	1.0 (13)	5.5 (34)	175 (90)
112,10-102,9	232 273.646	4.6	23 (18)	1.5 (19)	5.0 (39)	120 (90)
112,9-102,8	237 896.684	8.2	45 (9)	2.8 (3)	3.1 (15)	148 (42)
			HNCO			
40,4-30,3	87 925.237	3.2	162 (3)	4.0(1)	3.9 (1)	671 (11)
$5_{1.5} - 4_{1.4}$	109 495.996	5.4	38 (6)	2.0 (3)	4.5 (11)	182 (35)
$5_{0,5}-4_{0,4}$	109 905.749	5.9	254 (6)	3.9(1)	3.2 (1)	853 (18)
51,4-41,3	110 298.089	5.4	16 (5)	5.4 (6)	5.4 (18)	90 (23)
61,6-51,5	131 394.230	5.4	39 (17)	2.5 (8)	5.2 (27)	217 (88)
60,6-50,5	131 885.734	7.3	263 (8)	3.9(1)	3.0(1)	825 (18)
61,5-51,4	132 356.701	6.9	39 (8)	3.6 (5)	5.3 (13)	218 (48)
71,7-61,6	153 291.935	9.5	73 (27)	5.1 (5)	3.0 (16)	230 (90)
70,7-60,6	153 865.086	12.3	224 (18)	3.9 (1)	4.0 (3)	950 (50)
71,6-61,5	154 414.765	11.1	57 (28)	2.7 (7)	3.4 (21)	200 (90)
101,10-91,9	218 981.009	6.2	103 (8)	2.8 (2)	6.0 (5)	664 (47)
100,10-90,9	219 798.274	4.4	260 (12)	3.3 (1)	5.7 (3)	1580 (60)
10 _{1,9} –9 _{1,8}	220 584.751	6.4	85 (10)	2.9 (3)	6.2 (7)	560 (50)
11 _{1,11} -10 _{1,10}	240 875.727	14.5	132 (16)	2.8 (3)	6.6 (9)	930 (100)
$12_{0,12} - 11_{0,11}$	263 748.625	7.9	270 (70)	3.7 (6)	5.6 (15)	1640 (360)
$12_{1,11} - 11_{1,10}$	264 693.655	6.9	119 (11)	3.4 (2)	6.6 (6)	840 (60)

Note. ^w Transition weakly detected (see Table B1) but included in the analysis for completeness.

Table B10.	SVS13A: Gaussian fits to the detected NH ₂ CHO and HNCO lines.
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Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV (\rm mK \ km \ s^{-1})$
			NH ₂ CHO			
50,5-40,4	105 464.219	3.9	16(1)	8.1 (2)	1.8 (5)	14 (1)
71,7-61,6	142 701.325	6.4	28 (3)	8.6 (2)	2.2 (5)	22 (5)
73,4-63,3	148 709.018	7.1	26 (2)	7.1 (3)	2.2 (7)	22 (3)
102,9-92,8	211 328.960	7.3	62 (5)	7.5 (5)	4.4 (14)	31 (2)
103,8-93,7	212 572.837	7.2	43 (7)	8.4 (2)	2.7 (6)	29 (2)
$11_{4.8} - 10_{4.7}$	233 734.724	6.6	26 (2)	6.9 (4)	3.0 (9)	23 (5)
114.7-104.6	233 745.613	7.5	26 (2)	8.2 (3)	1.5 (7)	26(1)
113.9-103.8	233 896.577	6.6	49 (14)	8.2 (4)	2.9 (9)	20 (3)
112,9-102,8	237 896.684	6.9	48 (6)	8.1 (3)	4.3 (8)	28 (4)
$11_{1.10} - 10_{1.9}$	239 951.800	8.1	66 (7)	7.6 (4)	3.4 (9)	23 (2)
123,10-113,9	255 225.651	5.6	45 (2)	8.8 (12)	3.3 (36)	24 (4)
$12_{1,11} - 11_{1,10}$	261 327.450	8.7	26 (2)	8.4 (2)	4.6 (6)	38 (5)
131,13-121,12	263 542.236	7.6	54 (5)	7.7 (6)	4.4 (15)	32 (5)
			HNCO			
41,4-31,3	87 597.330	3.5	11 (3)	6.4 (4)	3.3 (8)	40 (9)
40,4-30,3	87 925.237	9.1	46 (4)	8.6 (2)	3.1 (6)	155 (23)
4 _{1,3} -3 _{1,2} ^w	88 239.020	3.0	13 (3)	7.3 (4)	3.3 (20)	45 (17)
51,5-41,4	109 495.996	3.9	16(1)	7.1 (2)	2.0 (5)	35 (8)
50,5-40,4*	109 905.749	5.4	76 (5)	8.5 (1)	1.2 (1)	100 (8)
51,4-41,3	110 298.089	6.2	28 (4)	8.2 (3)	4.7 (8)	139 (19)
61,6-51,5	131 394.230	6.6	19 (3)	8.2 (4)	3.6 (8)	72 (15)
60,6-50,5*	131 885.734	6.0	77 (10)	8.5 (1)	1.6 (3)	129 (15)
71,7-61,6	153 291.935	5.7	25 (2)	8.5 (5)	5.1 (13)	140 (30)
70,7-60,6*	153 865.086	6.1	70 (6)	7.5 (1)	3.4 (5)	258 (19)
71,6-61,5	154 414.765	6.0	32 (3)	8.3 (6)	4.4 (15)	149 (40)
$10_{1,10} - 9_{1,9}$	218 981.009	6.8	46 (3)	8.2 (2)	4.3 (4)	212 (15)
$10_{0,10} - 9_{0,9}$	219 798.274	7.1	89 (4)	8.4 (1)	3.2 (2)	309 (16)
10 _{1,9} -9 _{1,8}	220 584.751	6.0	35 (3)	8.5 (4)	3.7 (9)	136 (27)
111,11-101,10	240 875.727	6.7	46 (4)	8.2 (5)	4.9 (17)	236 (66)
111,10-101,9	242 639.704	8.7	42 (5)	8.1 (3)	4.6 (6)	206 (24)
121,12-111,11	262 769.477	9.9	68 (4)	8.5 (2)	5.3 (5)	380 (32)
120,12-110,11	263 748.625	9.3	60 (5)	8.2 (4)	3.4 (9)	213 (48)
121,11-111,10	264 693.655	8.9	39 (4)	8.5 (3)	4.4 (6)	183 (23)

Notes. * Transition affected by emission at OFF position: lower limit point in the rotational diagram.

 $^{\mathrm{w}}$ Transition weakly detected (see Table B1) but included in the analysis for completeness.

Table B11. Cep	E: Gaussian f	fits to the detected	d NH ₂ CHO and HNCO lines.
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Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\frac{\int T_{\rm mb} \mathrm{d}V}{(\mathrm{mK \ km \ s^{-1}})}$
			NH ₂ CHO)		
40.4-30.3	84 542.330	1.9	7.7 (1)	-11.5 (4)	2.8 (10)	23 (7)
42,2-32,1	85 093.272	1.3	4.3 (1)	-11.8 (3)	3.5 (7)	16 (3)
$4_{1,3}-3_{1,2}^{w}$	87 848.873	1.2	6.3 (1)	-10.5(2)	2.6 (5)	17 (3)
$5_{1.5} - 4_{1.4}$	102 064.267	1.6	7.6(1)	-10.6(2)	1.7 (5)	13 (3)
50,5-40,4	105 464.219	2.1	4.1 (1)	-10.6 (6)	2.9 (11)	13 (4)
			HNCO			
40.4-30.3	87 925.237	1.4	90 (4)	-11.1(1)	1.9(1)	179 (3)
$5_{0.5} - 4_{0.4}$	109 905.749	2.8	104 (12)	-11.1 (1)	2.4 (1)	262 (6)
60,6-50,5	131 885.734	7.8	128 (9)	-11.1 (1)	2.2 (3)	299 (23)
$10_{0,10} - 9_{0,9}$	219 798.274	7.5	45 (10)	-10.1 (73)	8.0 (80)	380 (65)
$12_{0,12} - 11_{0,11}$	263 748.625	6.0	38 (8)	-10.2 (5)	5.6 (14)	226 (45)

Note. ^w Transition weakly detected (see Table **B1**) but included in the analysis for completeness.

Table B12. OMC-2 FIR 4: Gaussian fits to the detected NH₂CHO and HNCO lines.

Transition	v (MHz)	rms (mK)	T _{peak} (mK)	$V_{\rm lsr}$ (km s ⁻¹)	ΔV (km s ⁻¹)	$\int T_{\rm mb} dV$ (mK km s ⁻¹)
	. ,	. ,	NH ₂ CHO			
40.4-30.3	84 542.330	2.8	15 (2)	10.8 (2)	2.2 (4)	35 (5)
41.3-31.2	87 848.873	2.8	9 (2)	11.0 (4)	3.6 (6)	37 (7)
$5_{1.5}-4_{1.4}$	102 064.267	3.5	15 (1)	11.0 (2)	2.9 (5)	46 (6)
$5_{0.5}-4_{0.4}$	105 464.219	4.7	16(1)	11.5 (4)	3.7 (10)	64 (15)
$5_{2,4}-4_{2,3}$	105 972.599	4.6	15 (5)	11.3 (2)	1.7 (5)	28 (8)
$6_{1,5}-5_{1,4}$	131 617.902	5.6	29 (3)	11.2 (2)	3.8 (6)	115 (13)
7 _{1,7} –6 _{1,6}	142 701.325	6.4	28 (3)	11.5 (2)	3.1 (7)	91 (14)
73.4-63.3	148 709.018	5.6	26 (2)	11.5 (2)	1.8 (5)	51 (10)
7 _{1,6} –6 _{1,5}	153 432.176	8.2	40 (3)	11.8 (3)	2.7 (10)	114 (31)
$10_{1,10} - 9_{1,9}$	203 335.761	10.8	52 (4)	11.3 (2)	2.8 (5)	156 (20)
100,10-90,9	207 679.189	8.8	49 (3)	11.6(1)	1.8 (4)	97 (14)
103.8-93.7	212 572.837	12.7	43 (7)	11.7 (3)	2.9 (6)	131 (23)
$11_{1,11} - 10_{1,10}$	223 452.512	12.6	44 (6)	11.7 (5)	3.0 (12)	142 (48)
$11_{0,11} - 10_{0,10}$	227 605.658	14.5	61 (10)	11.9 (3)	3.7 (7)	237 (33)
113.9-103.8	233 896.577	16.9	49 (14)	11.6 (3)	2.9 (10)	151 (36)
112.9-102.8	237 896.684	10.1	48 (6)	11.4 (2)	3.7 (6)	186 (23)
$11_{1,10} - 10_{1,9}$	239 951.800	10.5	66 (7)	11.5 (1)	2.5 (4)	175 (21)
$12_{2,11} - 11_{2,10}$	253 165.793	12.4	44 (9)	11.2 (3)	3.2 (7)	152 (27)
$12_{4.9} - 11_{4.8}^{W}$	255 058.533	12.5	39 (12)	11.6 (2)	1.9 (4)	80 (17)
123,10-113,9	255 225.651	11.4	45 (2)	11.5 (4)	1.6 (9)	74 (4)
123,9-113,8	255 871.830	11.8	41 (7)	11.8 (2)	2.7 (5)	119 (19)
			HNCO			
40,4-30,3	87 925.237	2.8	128 (13)	11.2(1)	2.5(1)	344 (6)
$4_{1,3}-3_{1,2}^{W}$	88 239.020	2.3	9 (2)	12.7 (3)	3.1 (6)	31 (5)
$5_{0.5} - 4_{0.4}$	109 905.749	6.9	224 (22)	11.4 (1)	2.1(1)	512 (14)
$5_{1.4} - 4_{1.3}^{W}$	110 298.089	6.2	18 (5)	12.8 (2)	1.4 (5)	26 (7)
60,6-50,5	131 885.734	4.7	267 (27)	11.2 (1)	2.7 (1)	775 (11)
70,7-60,6	153 865.086	8.8	304 (38)	11.2 (1)	3.0 (4)	962 (89)
100,10-90,9	219 798.274	9.2	343 (28)	11.4 (1)	2.5 (1)	918 (17)
101,9-91,8	220 584.751	11.0	36 (1)	11.7 (4)	2.1 (13)	79 (35)
$12_{0,12} - 11_{0,11}$	263 748.625	5.1	202 (9)	11.9 (3)	3.0 (5)	648 (24)

Note. ^w Transition weakly detected (see Table B1) but included in the analysis for completeness.

APPENDIX C: FIGURES

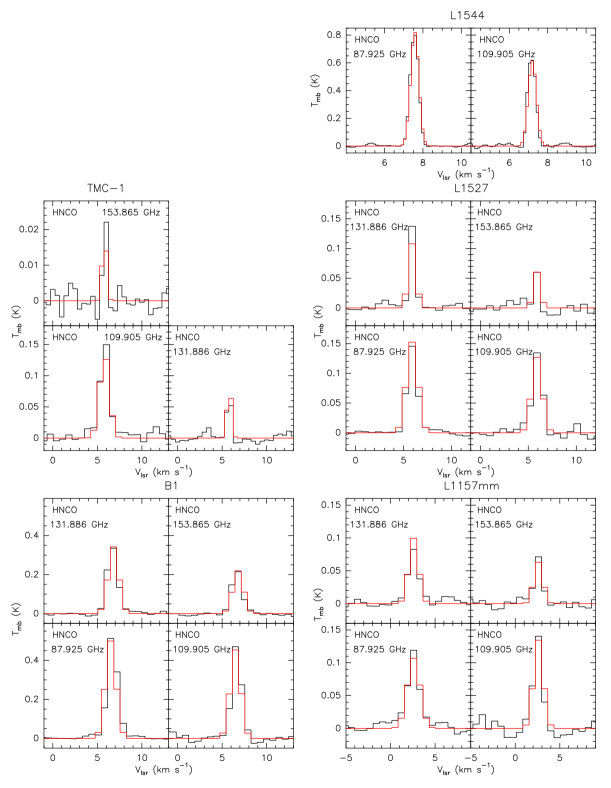


Figure C1. HNCO observed spectral lines (black) in L1544, TMC-1, B1, L1527, and L1157mm, and the spectra predicted by best-fitting LTE model (red).

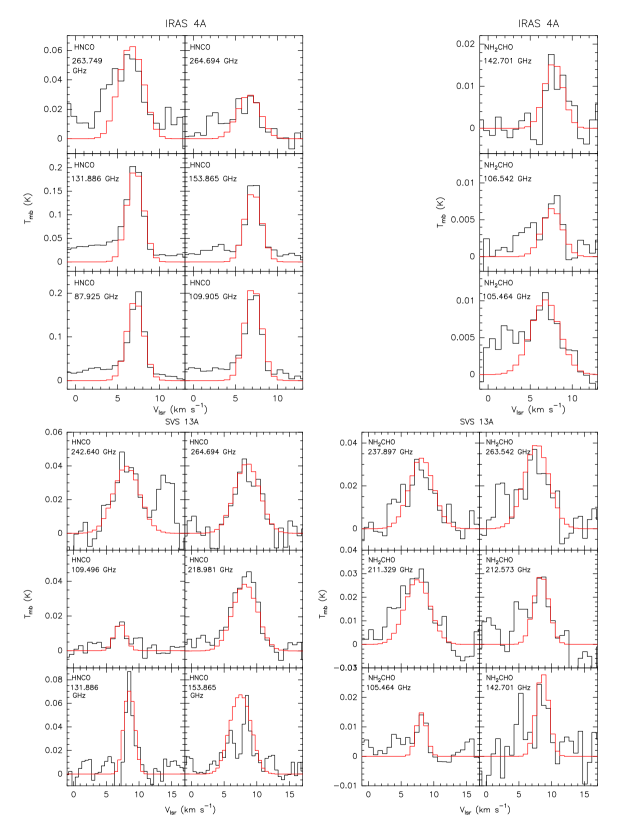


Figure C2. Sample of HNCO (left) and NH₂CHO (right) observed spectral lines (black) in IRAS 4A and SVS13A (compact solution), and the spectra predicted by best-fitting LTE model (red).

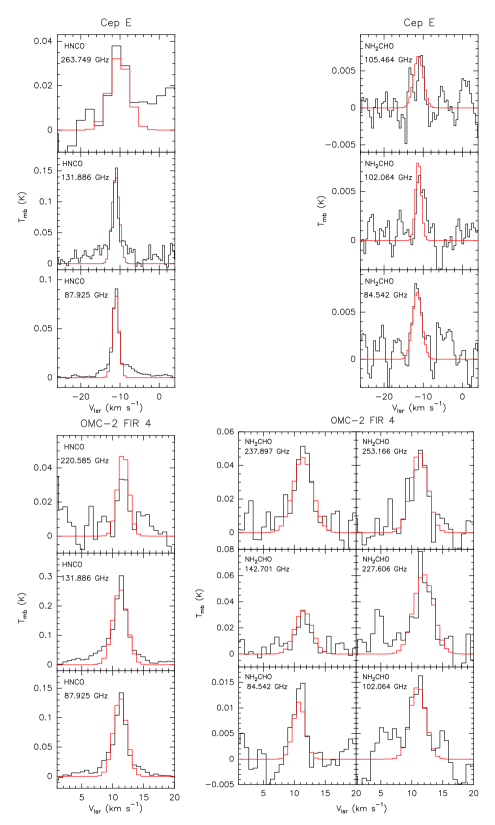


Figure C3. Sample of HNCO (left) and NH₂CHO (right) observed spectral lines (black) in Cep E and OMC-2 FIR 4 (extended solutions), and the spectra predicted by best-fitting LTE model (red).

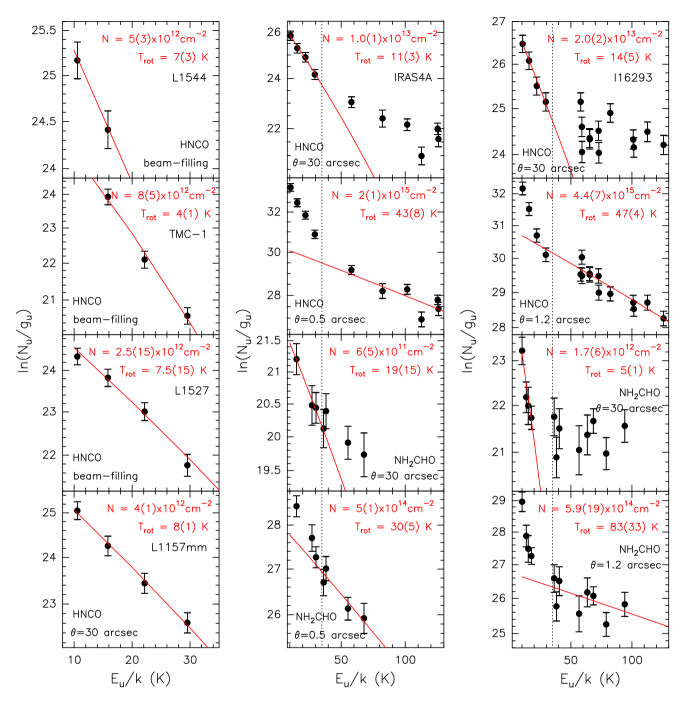


Figure C4. Rotational diagrams for L1544, TMC-1, L1527, and L1157mm (left), IRAS 4A (middle), and I16293 (right). Data points are depicted in black. The red lines correspond to the best fit to the data points. The dashed vertical lines in the middle and right panels indicate the upper-level energy (35 K) at which the division of the two-component fitting was made.

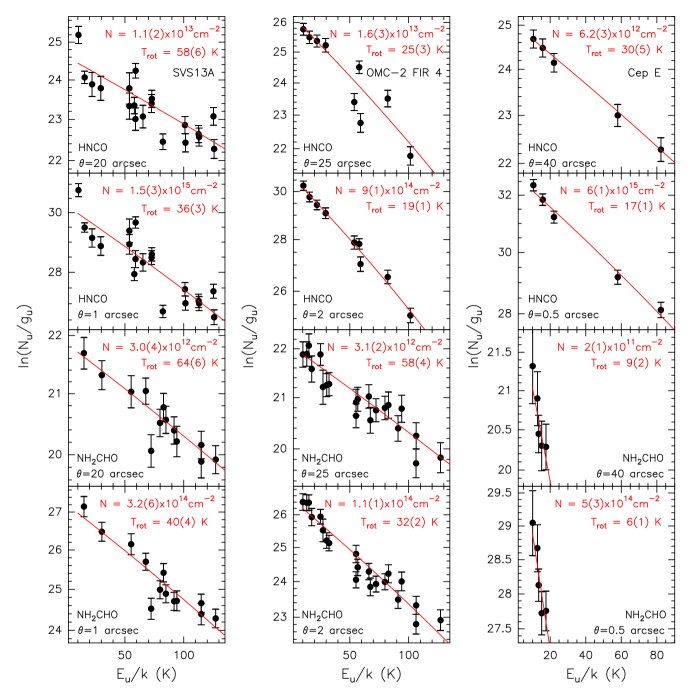


Figure C5. Rotational diagrams for SVS13A (left), OMC-2 FIR 4 (middle), and Cep E (right). Data points are depicted in black. The red lines correspond to the best fit to the data points.

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