LETTER TO THE EDITOR

The ALMA-PILS survey: First detections of deuterated formamide and deuterated isocyanic acid in the interstellar medium

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ABSTRACT

Formamide (NH2CHO) has previously been detected in several star-forming regions and is thought to be a precursor for different prebiotic molecules. Its formation mechanism is still debated, however. Observations of formamide, related species and their isopotologues may provide useful clues to the chemical pathways leading to their formation. The Protostellar Interferometric Line Survey (PILS) represents an unbiased high angular resolution and sensitivity spectral survey of the low-mass protostellar binary IRAS 16293– 2422 with the Atacama Large Millimeter/submillimeter Array (ALMA). We detect for the first time the three singly deuterated forms of NH2CHO (NH2CDO, cis- and trans-NHDCHO) as well as DNCO towards the component B of this binary source. The images reveal that the different isotopologues all are present in the same region. Based on the observations of the ¹³C isotopologues of formamide and a standard $^{12}C/$ mamide and a standard ${}^{12}C/{}^{13}C$ ratio, the deuterium fractionation is found to be similar for the three different forms with a value of about 2%. The DNCO/HNCO ratio is also comparable to the D/H ratio of formamide (∼1%). These results are in agreement with the hypothesis that NH₂CHO and HNCO are chemically related through grain surface formation.

Key words. astrochemistry – astrobiology – stars: formation – stars: protostars – ISM: molecules – ISM: individual object (IRAS 16293–2422)

1. Introduction

Formamide (NH2CHO), also known as methanamide, contains the amide bond $(-N-C(=O))$, which plays an important role in the synthesis of proteins. This molecule is a precursor for potential compounds of genetic and metabolic interest [\(Saladino et al.](#page-3-0) [2012\)](#page-3-0). Interestingly, it is present in various astrophysical environments: high-mass star-forming regions (e.g, [Bisschop et al.](#page-3-1) [2007;](#page-3-1) [Adande et al. 2013\)](#page-3-2), low-mass protostars [\(Kahane et al.](#page-3-3) 2007; Adande et al. 2013), low-mass protostars (Kahane et al. [2013;](#page-3-3) [López-Sepulcre et al. 2015\)](#page-3-4), shocked regions [\(Yamaguchi](#page-3-5) [et al. 2012;](#page-3-5) [Mendoza et al. 2014\)](#page-3-6), a translucent cloud [\(Corby](#page-3-7) [et al. 2015\)](#page-3-7), comets [\(Bockelée-Morvan et al. 2000;](#page-3-8) [Biver et al.](#page-3-9) [2014;](#page-3-9) [Goesmann et al. 2015\)](#page-3-10) and even an extragalactic source [\(Muller et al. 2013\)](#page-3-11).

The formation of formamide is still not clearly understood: several routes have been proposed, both in the gas phase and on the grain surfaces. In the gas phase, many ion-molecule reactions have been ruled out as not sufficiently efficient due to endothermicity or high energy barriers (see e.g. [Redondo et al.](#page-3-12) [2014a,](#page-3-12)[b\)](#page-3-13). A neutral-neutral reaction between H_2CO and $NH₂$

was however shown to be barrierless and could account for the abundance of formamide in some sources [\(Barone et al. 2015\)](#page-3-14). On the grain surface, formamide is suggested to form through the reaction between HCO and NH² [\(Jones et al. 2011;](#page-3-15) [Garrod](#page-3-16) [2013\)](#page-3-16) and/or hydrogenation of isocyanic acid, HNCO. In particular, the latter suggestion is supported by a strong correlation between the HNCO and NH₂CHO abundances in different sources [\(Bisschop et al. 2007;](#page-3-1) [Mendoza et al. 2014;](#page-3-6) [López-Sepulcre et al.](#page-3-4) [2015\)](#page-3-4). However, an experiment based on the H bombardment of HNCO at low temperature has recently shown that this reaction is not efficient in cold environments [\(Noble et al. 2015\)](#page-3-17). Instead, other pathways to HNCO and $NH₂CHO$ on grains have been suggested, either with or without UV or ion bombardment (see e.g. Kaňuchová et al. 2016 and references therein).

Measurements of isotopic fractionation may help to constrain formation pathways of molecules as isotopic fractionation (especially deuteration) is sensitive to physical conditions such as density and temperature. Until recently, the study of deuteration in solar-type protostars was mainly limited to relatively small and abundant molecules, such as H_2O , HCO^+ ,

HCN, H_2CO , and CH₃OH. Even though the deuterium fractionation is known to be enhanced in low-mass protostars (see e.g., [Ceccarelli et al. 2007\)](#page-3-19), measurements of lines of deuterated complex organic molecules (COMs) still require high sensitivity observations. So far, only deuterated methyl formate and dimethyl ether have been detected towards the low-mass protostar IRAS 16293–2422 (hereafter IRAS16293) by [Demyk](#page-3-20) [et al.](#page-3-20) [\(2010\)](#page-3-20) and [Richard et al.](#page-3-21) [\(2013\)](#page-3-21). With the Atacama Large Millimeter/submillimeter Array (ALMA), it is now possible to search for the isotopologues of complex and less abundant species. In this Letter, we report the first detection of the three singly deuterated forms of formamide as well as DNCO towards IRAS16293. These observations mark the first detections of those isotopologues in the interstellar medium.

2. Observations

An ALMA unbiased spectral survey of the binary protostar IRAS16293 was recently carried out in the framework of the "Protostellar Interferometric Line Survey"^{[1](#page-1-0)} (PILS; [Jørgensen](#page-3-22) [et al. submitted\)](#page-3-22). The observations were centered on a position at equal distance between the sources A and B that are separated by ∼5 ⁰⁰. A full description of the survey and the data reduction can be found in [Jørgensen et al.](#page-3-22) [\(submitted\)](#page-3-22). For this work, we use the part of the large spectral survey obtained in Band 7 between 329.15 GHz and 362.90 GHz both with the 12m array and the Atacama Compact Array (ACA). The spectral resolution of these observations is 0.244 MHz (i.e. ~0.2 km s⁻¹). After combination of the 12m and ACA data, the final spectral line datacubes show a sensitivity better than 5 mJy beam⁻¹ km s⁻¹. The beam sizes range between 0.4 " and 0.7 ". Additional observations in Bands 3 and 6 cover narrow spectral ranges and consequently a very limited number of transitions of formamide isotopologues. After the analysis of Band 7, we checked that the results are consistent with these lower frequency observations.

3. Analysis and results

To search for the isotopologues of formamide, we use the spectrum extracted at the same position as in [Lykke et al.](#page-3-23) [\(to be sub](#page-3-23)[mitted\)](#page-3-23), i.e. a position offset by ~ 0.5 ^o from the continuum peak of source B in the South West direction $(\alpha_{12000}=16^{h}32^{m}22558,$
 $\delta_{12000}=24^{\circ}28'32.8'')$ Although the lines are brighter at the po- δ_{J2000} =-24°28′32.8″). Although the lines are brighter at the position of the continuum peak, the presence of both absorption and emission makes analysis difficult. At the selected position, most of the lines present Gaussian profiles and are relatively bright compared to other positions. In source A, the lines are quite broad leading to significant line confusion that prevents the search for isotopologues of complex species (e.g. [Jørgensen](#page-3-24) [et al. 2012\)](#page-3-24). This Letter is therefore focused on source B only.

We identify several unblended lines that can be assigned to the three singly deuterated forms of $NH₂CHO$ and to $NH₂$ ¹³CHO, DNCO, and HN¹³CO (see Table [1\)](#page-1-1). These mark the first detections of NH2CDO, cis-NHDCHO, trans-NHDCHO and DNCO in the interstellar medium. The list of unblended lines can be found in the Appendix. Maps of the integrated line emission from representative lines from the different isotopologues towards source B are shown in Figure [1.](#page-2-0) The emission of the different lines clearly arise from a similar compact region in the vicinity of IRAS16293B. A hole is observed in the maps due to the absorptions that are produced against the strong continuum at the continuum peak position. For DNCO the larger

Table 1. Number of lines used in the analysis of the isotopologues of $NH₂CHO$ and HNCO and column densities derived for $T_{ex} = 300$ K and a source size of 0.5".

Species	# of lines	$E_{\rm up}$ (K)	N (cm ⁻²)
$NH2$ CDO	12	$146 - 366$	2.1×10^{14}
cis-NHDCHO	11	$146 - 307$	2.1×10^{14}
trans-NHDCHO	11	$151 - 332$	1.8×10^{14}
NH ₂ ¹³ CHO	10	$152 - 428$	1.5×10^{14}
15 NH ₂ CHO			$\leq 1.0\times 10^{14}$ $^{(a)}$
NH ₂ CH ¹⁸ O			$\leq 0.8 \times 10^{14}$ $^{(a)}$
DNCO	4	$150 - 751$	3.0×10^{14}
HN ¹³ CO	8	$127 - 532$	4.0×10^{14}
$H^{15}NCO$			$\leq 2.0\times 10^{14}$ $^{(a)}$
HNC ¹⁸ O			$\leq 1.5 \times 10^{14}$ $^{(a)}$

Notes. ^(a) 3σ upper limit.

beam size for the observations of this transition masks the absorption. The spatial variations that are observed among the different species are probably due to different line excitation or line brightness. In particular, HNCO seems to be slightly more extended than $NH₂CHO$, but this is most likely due to the fact that the HNCO lines are particularly bright compared to the HNCO and formamide isotopologues.

To constrain the excitation temperatures and column densities of the different species, we produce a grid of synthetic spectra assuming Local Thermodynamical Equilibrium (LTE). We predict the spectra for different excitation temperatures between 100 and 300 K with a step of 25 K and for different column densities between 1×10^{13} and 1×10^{17} cm⁻². First, the column density is roughly estimated using relatively large steps, then refined using smaller steps around the best fit solution. We determine the best fit model using a χ^2 method comparing the observed
and synthetic spectra at +0.5 MHz around the rest frequency of and synthetic spectra at ± 0.5 MHz around the rest frequency of the predicted emission lines. We carefully check that the best fit model does not predict any lines not observed in the spectra. For the deuterated forms, the models are in agreement with the observations for excitation temperatures between 100 and 300 K. However, for $NH₂$ ¹³CHO and HN¹³CO, a model with a high excitation temperature accounts much better for the observed emission than a model with a low excitation temperature (see Figs [B.4](#page-8-0) and [B.6\)](#page-8-1). An excitation temperature of 300 K was consequently adopted for the analysis of the different isotopologues. This excitation temperature is similar to that derived for glycolaldehyde and ethylene glycol [\(Jørgensen et al. 2012,](#page-3-24) submitted), but higher than what is found for acetaldehyde, ethylene oxide and propanal (∼125 K, [Lykke et al. to be submitted\)](#page-3-23). The derived column densities, assuming a linewidth of 1 km s[−]¹ and a source size of 0.5" [\(Jørgensen et al. submitted;](#page-3-22) [Lykke et al. to](#page-3-23) [be submitted\)](#page-3-23), are summarized in Table [1.](#page-1-1) The uncertainties on the column densities are all estimated to be within a factor of 2 (including the uncertainty on both the excitation temperature and the baseline subtraction). The upper limits are estimated visually by comparison of the synthetic spectra with the observations on the entire spectral range. Figure [2](#page-2-1) shows three lines of each isotopologue with the best-fit model. The models for all the lines are shown in Appendix B.

The column densities of $NH₂^{13}CHO$ and $HN¹³CO$ are estimated to be 1.5×10^{14} cm⁻² and 4×10^{14} cm⁻², respectively. Assuming a ${}^{12}C/{}^{13}C$ ratio of 68 [\(Milam et al. 2005\)](#page-3-25), the column densities for the main isotopologues of formamide and isocyanic acid are predicted to be 1×10^{16} cm⁻² and 3×10^{16} cm⁻².

¹ http://youngstars.nbi.dk/PILS/

Fig. 1. Integrated intensity maps of NH₂CHO, HNCO and their isotopologues towards source B. The position of the continuum peak of source B is indicated with a red cross, while the position where the spectrum was extracted is shown with a red circle. The beam sizes are shown in grey in the bottom right corner of each panel. The contour levels start for the main isotopologue of HNCO at 0.05 Jy km s[−]¹ with a step of 0.05 Jy km s⁻¹. For the other species, the levels are 0.02, 0.03, 0.04, 0.06, 0.08, 0.1 and 0.12 Jy km s⁻¹.

With these column densities, several $NH₂CHO$ lines and all of the HNCO lines are overproduced, indicating that they are optically thick. The model of formamide is, however, in agreement with the few lines with the lowest opacities (see Figs. [B.7](#page-9-0) and [B.8\)](#page-9-1). $NH₂CH¹⁸O$ has also been searched for, but is not detected with a 3 σ upper limit of 8×10^{13} cm⁻². The non-detection of this isotopologue is consistent with the ¹⁶O/¹⁸O ratio of 560 in the inisotopologue is consistent with the $\rm ^{16}O/^{18}O$ ratio of 560 in the in-terstellar medium [\(Wilson 1999\)](#page-3-26), which gives $N(NH_2CH^{18}O)$ = 2×10^{13} cm⁻². Similarly, HNC¹⁸O is not detected either with a 3σ upper limit of 1.5×10^{14} cm⁻², which is consistent with its expected column density of 5×10^{13} cm⁻² expected column density of 5×10^{13} cm⁻².

Using the column densities derived for the 13 C isotopologues and a standard ${}^{12}C/{}^{13}C$ ratio, the deuterium fractionation in $NH₂CHO$ is about 2% for the three deuterated forms and the DNCO/HNCO ratio is similar (~1%). If the ¹²C/¹³C ratio is lower (∼30) as reported for glycolaldehyde by [Jørgensen](#page-3-22) [et al.](#page-3-22) [\(submitted\)](#page-3-22), the D/H ratios of formamide and isocyanic acid would be about 4-5% and 2-3%, respectively.

Fig. 2. *Black*: Detected lines of NH₂CDO, cis-NHDCHO, trans-NHDCHO, NH² ¹³CHO, DNCO and HN¹³CO. *Red*: Best-fit model.

We also search for the $15N$ isotopologues of formamide and isocyanic acid. A couple of transitions could tentatively be assigned to $\rm ^{15}NH_2CHO$, but these lines are close to the noise level and possibly blended with other species. For $H^{15}NCO$, the uncertainties on the frequencies of some of the transitions are rather large, preventing any firm detection. Based on a standard $^{12}C/^{13}C$ ratio, lower limits of 100 and 138 are obtained for the $^{14}N/^{15}N$ ratios of formamide and HNCO respectively.

4. Discussion and conclusion

Our derived ratio in IRAS16293 for HNCO/NH2CHO, ∼3, is consistent with the ratios found in warm sources in previous studies [\(Bisschop et al. 2007;](#page-3-1) [Mendoza et al. 2014;](#page-3-6) [López-](#page-3-4)[Sepulcre et al. 2015\)](#page-3-4). Thanks to our interferometric observations, we also confirm that these two species are spatially correlated. The deuterium fractionation ratios of these two molecules are also similar, reinforcing the hypothesis that they are chemically related. We discuss here possible scenarios for the formation of these species in the warm inner regions of protostars.

Assuming that the deuteration of formaldehyde in the region probed by the ALMA observations of formamide is similar to the value derived with single-dish observations (∼15%, [Loinard](#page-3-27) [et al. 2000\)](#page-3-27), we can discuss the possibility for the gas-phase for-

mation mechanism proposed by [Barone et al.](#page-3-14) [\(2015\)](#page-3-14), H_2CO + $NH₂ \rightarrow NH₂CHO + H$. According to this reaction, the deuterated form NHDCHO would result from the reaction between NHD and H_2CO , while NH₂CDO would form from NH₂ and HDCO. We would consequently expect a higher deuteration for $NH₂CDO$ compared to the observations unless the reaction between $NH₂$ and HDCO leads more efficiently to $NH₂CHO$ and D compared to NH2CDO and H. Theoretical or experimental studies of the branching ratios of these reactions would be needed to rule out this scenario. The determination of the $HDCO/H₂CO$ ratio from the PILS survey is also necessary. Nevertheless, it should be noted that so far there is no proposed scenario in the gas phase that could explain the correlation with HNCO.

Although it was recently shown that $NH₂CHO$ does not form by hydrogenation of HNCO on grain surfaces [\(Noble et al.](#page-3-17) [2015\)](#page-3-17), several other proposed mechanisms exist in the literature. Both species can be formed through barrierless reactions in ices through NH + CO \rightarrow HNCO and NH₂ + H₂CO \rightarrow NH₂CHO + H, as demonstrated experimentally [\(Fedoseev et al. 2015,](#page-3-28) [2016\)](#page-3-29). Alternatively, both species are formed through ion bombardment of $H_2O:CH_4:N_2$ mixtures (Kaňuchová et al. 2016) or UV irradiation of CO:NH3:CH3OH and/or HNCO mixtures (e.g. [Demyk](#page-3-30) [et al. 1998;](#page-3-30) [Raunier et al. 2004;](#page-3-31) [Jones et al. 2011;](#page-3-15) [Henderson](#page-3-32) [& Gudipati 2015\)](#page-3-32). Quantitative gas-grain modeling under conditions representative of IRAS16293 are needed to assess which of these grain surface routes dominates.

Ultimately, the HNCO and NH₂CHO deuterium fractionation level and pattern may also hold a clue to their formation routes. A particularly interesting result is that the three singly deuterated forms of formamide are found with similar abundances in IRAS16293. Contrary to the -CH functional group that is not affected by hydrogen isotope exchanges, the hydroxyl (-OH) and amine (-NH) groups are expected to establish hydrogen bonds and equilibrate with water [\(Faure et al.](#page-3-33) [2015\)](#page-3-33). This mechanism was proposed to explain the different CH3OD/CH3OH (∼1.8%) and CH2DOH/CH3OH (∼37%) ratios derived in IRAS16293 [\(Parise et al. 2006\)](#page-3-34), as the water deuterium fractionation of water in the upper layers of the grain mantles where complex organic molecules form is about a few percent [\(Coutens et al. 2012,](#page-3-35) [2013;](#page-3-36) [Furuya et al. 2016\)](#page-3-37). We do not see such differences for formamide, for which all forms show a deuterium fractionation similar to the $CH₃OD/CH₃OH$ ratio and water. The deuterium fractionation of methanol from the PILS data needs to be investigated to know if the different deuterium fractionation ratios of the -CH and -OH groups are also observed at small scales.

In conclusion, we present in this Letter the first detection of the three singly deuterated forms of formamide and DNCO. The similar deuteration of these species and their similar spatial distributions favours the formation of these two species on grain surfaces. Further studies are, however, needed to rule out gas phase routes. These detections illustrate the strength of ALMA, and large spectral surveys such as PILS in particular, for the detections of deuterated complex molecules. Determinations of the deuterium fractionation for more complex molecules will help to constrain their formation pathways. The search for deuterated formamide in more sources is needed to reveal how variable the deuteration of formamide is, and if the similarity of the abundances of the three deuterated forms is common.

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Appendix A: Spectroscopic data

A list of unblended and optically thin lines used in the analysis is presented in Table [A.1.](#page-5-0) The spectroscopic data for $NH₂CHO$ $v=0$, NH₂CHO $v_{12}=1$, NH₂¹³CHO, ¹⁵NH₂CHO, NH₂CH¹⁸O, NH2CDO, cis-NHDCHO, trans-NHDCHO [\(Kurland & Bright](#page-3-38) [Wilson 1957;](#page-3-38) [Kukolich & Nelson 1971;](#page-3-39) [Hirota et al. 1974;](#page-3-40) [Gardner et al. 1980;](#page-3-41) [Moskienko & Dyubko 1991;](#page-3-42) [Vorob'eva &](#page-3-43) [Dyubko 1994;](#page-3-43) [Blanco et al. 2006;](#page-3-44) [Kryvda et al. 2009;](#page-3-45) [Motiyenko](#page-3-46) [et al. 2012;](#page-3-46) [Kutsenko et al. 2013\)](#page-3-47) and HNCO [\(Kukolich & Nel](#page-3-39)[son 1971;](#page-3-39) [Hocking et al. 1975;](#page-3-48) [Niedenho](#page-3-49)ff et al. [1995;](#page-3-49) [Lapinov](#page-3-50) [et al. 2007\)](#page-3-50) come from the CDMS database [\(Müller et al.](#page-3-51) [2001,](#page-3-51) [2005\)](#page-3-52), while the data for DNCO, $HN^{13}CO$, $H^{15}NCO$ and HNC18O [\(Hocking et al. 1975\)](#page-3-48) are taken from the JPL database [\(Pickett et al. 1998\)](#page-3-53). It should be noted that there are significant differences for the predicted frequencies of the main isotopologue of $NH₂CHO$ between CDMS and JPL (> 1 MHz). A better agreement is found with the observations for the most recent entry in CDMS. For some of the HNCO isotopologues, there is a lack of published spectroscopic data at high frequencies. In particular for $H^{15}NCO$, the range of uncertainty for some of the frequencies is quite high. As the HN13CO transitions appeared all slightly shifted compared to the observations, we applied a correction of ⁺0.5 MHz to model the lines.

The column densities of the formamide isotopologues given in Table [1](#page-1-1) were corrected by a factor of 1.5 to take into account the contribution of the vibrational states for an excitation temperature of 300 K.

Species	Transition	Frequency	$E_{\rm up}$	A_{ij}	$g_{\rm up}$
		(MHz)	(K)	(s^{-1})	
$NH2$ CDO	$(17017 - 16016)$	329995.2	145.6	$2.\overline{64 \times 10^{-3}}$	105
$NH2$ CDO	$(1697 - 1596)$	333363.6	308.9	1.87×10^{-3}	99
NH ₂ CDO	$(1698 - 1597)$	333363.6	308.9	1.87×10^{-3}	99
$NH2$ CDO	$(16710 - 1579)$	333696.6	240.7	2.22×10^{-3}	99
$NH2$ CDO	$(1679 - 1578)$	333696.6	240.7	2.22×10^{-3}	99
$NH2$ CDO	$(16413 - 15412)$	335234.9	170.5	2.61×10^{-3}	99
$NH2$ CDO	$(16313 - 15312)$	342320.7	156.9	2.86×10^{-3}	99
$NH2$ CDO	$(17116 - 16115)$	351988.3	158.1	3.18×10^{-3}	105
$NH2$ CDO	$(17107 - 16106)$	354151.5	366.4	2.15×10^{-3}	105
$NH2$ CDO	$(17108 - 16107)$	354151.5	366.4	2.15×10^{-3}	105
$NH2$ CDO	$(1798 - 1697)$	354257.0	325.9	2.37×10^{-3}	105
$NH2$ CDO	$(1799 - 1698)$	354257.0	325.9	2.37×10^{-3}	105
$NH2$ CDO	$(17810 - 1689)$	354416.0	289.6	2.56×10^{-3}	105
$NH2$ CDO	$(1789 - 1688)$	354416.0	289.6	2.56×10^{-3}	105
$NH2$ CDO	$(17711 - 16710)$	354661.3	257.7	2.74×10^{-3}	105
$NH2$ CDO	$(17710 - 1679)$	354661.3	257.7	2.74×10^{-3}	105
$NH2$ CDO	$(17512 - 16511)$	355800.2	206.7	3.04×10^{-3}	105
$NH2$ CDO	$(17413 - 16412)$	357938.5	187.8	3.20×10^{-3}	105
cis-NHDCHO	$(16313 - 15312)$	331372.8	156.0	2.59×10^{-3}	99
cis-NHDCHO	$(16214 - 15213)$	337248.5	146.0	2.79×10^{-3}	99
cis-NHDCHO	$(17216 - 16215)$	340520.3	158.0	2.87×10^{-3}	105
cis-NHDCHO	$(18118 - 17117)$	344878.9	160.8	3.02×10^{-3}	111
cis-NHDCHO	$(17810 - 1689)$	346444.0	306.6	2.39×10^{-3}	105
cis-NHDCHO	$(1789 - 1688)$	346444.0	306.6	2.39×10^{-3}	105
cis-NHDCHO	$(17711 - 16710)$	346586.8	269.8	2.56×10^{-3}	105
cis-NHDCHO	$(17710 - 1679)$	346586.8	269.8	2.56×10^{-3}	105
cis-NHDCHO	$(17612 - 16611)$	346826.8	238.0	2.70×10^{-3}	105
cis-NHDCHO	$(17611 - 16610)$	346827.5	238.0	2.70×10^{-3}	105
cis-NHDCHO	$(17315 - 16314)$	347115.8	172.0	2.99×10^{-3}	105
cis-NHDCHO	$(17512 - 16511)$	347268.9	211.1 189.2	2.83×10^{-3} 2.94×10^{-3}	105 105
cis-NHDCHO cis-NHDCHO	$(17414 - 16413)$ $(17314 - 16313)$	347827.8	173.0	3.15×10^{-3}	105
trans-NHDCHO	$(1789 - 1688)$	353047.5 333628.6	332.4	2.14×10^{-3}	105
trans-NHDCHO	$(17810 - 1689)$	333628.6	332.4	2.14×10^{-3}	105
trans-NHDCHO	$(17711 - 16710)$	333694.1	288.3	2.28×10^{-3}	105
trans-NHDCHO	$(17710 - 1679)$	333694.1	288.3	2.28×10^{-3}	105
trans-NHDCHO	$(17612 - 16611)$	333812.6	250.1	2.41×10^{-3}	105
trans-NHDCHO	$(17611 - 16610)$	333812.7	250.1	2.41×10^{-3}	105
trans-NHDCHO	$(17414 - 16413)$	334403.2	191.4	2.61×10^{-3}	105
trans-NHDCHO	$(18118 - 17117)$	336945.3	157.3	2.82×10^{-3}	111
trans-NHDCHO	$(18018 - 17017)$	338818.4	156.9	2.87×10^{-3}	111
trans-NHDCHO	$(17116 - 16115)$	338878.8	150.6	2.86×10^{-3}	105
trans-NHDCHO	$(18712 - 17711)$	353355.8	305.2	2.77×10^{-3}	111
trans-NHDCHO	$(18711 - 17710)$	353355.8	305.2	2.77×10^{-3}	111
trans-NHDCHO	$(18514 - 17513)$	353758.4	234.7	3.02×10^{-3}	111
trans-NHDCHO	$(18316 - 17315)$	354028.8	187.8	3.19×10^{-3}	111
trans-NHDCHO	$(18415 - 17414)$	354185.9	208.4	3.13×10^{-3}	111
NH ₂ ¹³ CHO	$(16106 - 15105)$	339170.1	427.9	1.75×10^{-3}	33
NH ₂ ¹³ CHO	$(16107 - 15106)$	339170.1	427.9	1.75×10^{-3}	33
NH ₂ ¹³ CHO	$(1697 - 1596)$	339179.6	373.0	1.97×10^{-3}	33
NH ₂ ¹³ CHO	$(1698 - 1597)$	339179.6	373.0	1.97×10^{-3}	33
NH ₂ ¹³ CHO	$(1688 - 1587)$	339213.5	323.8	2.16×10^{-3}	33
NH ₂ ¹³ CHO	$(1689 - 1588)$	339213.5	323.8	2.16×10^{-3}	33
NH ₂ ¹³ CHO	$(16 5 11 - 15 5 10)$	339672.1	210.9	2.61×10^{-3}	33
$NH2$ ¹³ CHO	$(16413 - 15412)$	340090.4	184.9	2.72×10^{-3}	33

Table A.1. Detected lines of $NH₂CHO$, HNCO and their isotopologues used in the analysis*(a)* .

Species	Transition	Frequency	$E_{\rm up}$	A_{ij}	$g_{\rm up}$
		(MHz)	(K)	(s^{-1})	
$\overline{\text{NH}_2{}^{13}\text{CHO}}$	$(16412 - 15411)$	340273.4	184.9	2.73×10^{-3}	33
NH ₂ ¹³ CHO	$(17117 - 16116)$	342156.0	151.5	2.95×10^{-3}	35
NH ₂ ¹³ CHO	$(1798 - 1697)$	360396.3	390.3	2.49×10^{-3}	35
NH ₂ ¹³ CHO	$(1799 - 1698)$	360396.3	390.3	2.49×10^{-3}	35
NH ₂ ¹³ CHO	$(17711 - 16710)$	360531.8	297.7	2.88×10^{-3}	35
NH ₂ ¹³ CHO	$(17710 - 1679)$	360531.8	297.7	2.88×10^{-3}	35
$NH2$ ¹³ CHO	$(18118 - 17117)$	361904.8	168.9	3.49×10^{-3}	37
$NH2CHO v=0$	$(16314 - 16215)$	331685.9	165.6	7.87×10^{-5}	$\overline{33}$
				5.49×10^{-5}	
$NH2CHO v=0$	$(827 - 716)$	334483.5	48.5		17
$NH2CHO v=0$	$(17315 - 17216)$	336733.0	183.0	8.2×10^{-5}	35
$NH2CHO v=0$	$(34331 - 34232)$	342029.5	645.9	1.07×10^{-4}	69
$NH2CHO v=0$	$(18316 - 18217)$	342511.1	201.3	8.57×10^{-5}	37
$NH2CHO v=0$	$(28424 - 28325)$	344545.8	464.1	1.15×10^{-4}	57
$NH2CHO v=0$	$(19317 - 19218)$	349051.7	220.7	8.99×10^{-5}	39
$NH2CHO v=0$	$(20318 - 20219)$	356379.8	241.1	9.47×10^{-5}	41
$NH2CHO v=0$	$(20119 - 19218)$	359119.4	221.2	8.45×10^{-5}	41
$NH2CHO v12=1$	$(17143 - 16142)$	360717.7	1144.3	1.12×10^{-3}	$\overline{35}$
$NH2CHO v12=1$	$(17144 - 16143)$	360717.7	1144.3	1.12×10^{-3}	35
DNCO	$(1711718 - 1611617)$	344629.4	172.9	5.92×10^{-4}	$\overline{37}$
DNCO	$(1711717 - 1611616)$	344629.4	172.9	5.90×10^{-4}	35
DNCO	$(1711716 - 1611615)$	344629.4	172.9	5.90×10^{-4}	33
DNCO	$(1701718 - 1601617)$			6.04×10^{-4}	37
		346556.2	149.7		
DNCO	$(1701717 - 1601616)$	346556.2	149.7	6.02×10^{-4}	35
DNCO	$(1701716 - 1601615)$	346556.2	149.7	6.02×10^{-4}	33
DNCO	$(17 5 12 18 - 16 5 11 17)$	346714.9	750.6	5.53×10^{-4}	37
DNCO	$(17 5 13 18 - 16 5 12 17)$	346714.9	750.6	5.53×10^{-4}	37
DNCO	$(17 5 13 16 - 16 5 12 15)$	346714.9	750.6	5.50×10^{-4}	33
DNCO	$(17 5 12 16 - 16 5 11 15)$	346714.9	750.6	5.50×10^{-4}	33
DNCO	$(17 5 13 17 - 16 5 12 16)$	346714.9	750.6	5.51×10^{-4}	35
DNCO	$(17 5 12 17 - 16 5 11 16)$	346714.9	750.6	5.51×10^{-4}	35
DNCO	$(1711618 - 1611517)$	348599.7	174.6	6.13×10^{-4}	37
DNCO	$(1711617 - 1611516)$	348599.7	174.6	6.10×10^{-4}	35
DNCO	$(1711616 - 1611515)$	348599.7	174.6	6.10×10^{-4}	33
HN ¹³ CO	$(1521316 - 1421215)$	329594.5	299.2	5.08×10^{-4}	$\overline{33}$
HN ¹³ CO	$(1521314 - 1421213)$	329594.5	299.2	5.06×10^{-4}	29
HN ¹³ CO	$(1521315 - 1421214)$	329594.5	299.2	5.06×10^{-4}	31
HN ¹³ CO	$(1501516 - 1401415)$	329673.4	126.6	5.18×10^{-4}	33
HN ¹³ CO	$(1501515 - 1401414)$	329673.4	126.6	5.16×10^{-4}	31
HN ¹³ CO	$(1501514 - 1401413)$	329673.4	126.6	5.15×10^{-4}	29
HN ¹³ CO	$(1511416 - 1411315)$			5.21×10^{-4}	33
		330860.2	170.2		
HN ¹³ CO	$(1511414 - 1411313)$	330860.2	170.2	5.19×10^{-4}	29
HN ¹³ CO	$(1511415 - 1411314)$	330860.2	170.2	5.19×10^{-4}	31
HN ¹³ CO	$(1611617 - 1511516)$	350340.3	186.1	6.20×10^{-4}	35
HN ¹³ CO	$(1611616 - 1511515)$	350340.3	186.1	6.18×10^{-4}	33
HN ¹³ CO	$(1611615 - 1511514)$	350340.3	186.1	6.18×10^{-4}	31
HN ¹³ CO	$(1631417 - 1531316)$	351427.6	531.9	6.07×10^{-4}	35
HN ¹³ CO	$(1631415 - 1531314)$	351427.6	531.9	6.04×10^{-4}	31
HN ¹³ CO	$(1631416 - 1531315)$	351427.7	531.9	6.04×10^{-4}	33
HN ¹³ CO	$(1631317 - 1531216)$	351427.7	531.9	6.07×10^{-4}	35
HN ¹³ CO	$(1631315 - 1531214)$	351427.7	531.9	6.04×10^{-4}	31
HN ¹³ CO	$(1631316 - 1531215)$	351427.7	531.9	6.04×10^{-4}	33
HN ¹³ CO	$(1621517 - 1521416)$	351548.3	316.1	6.19×10^{-4}	35
HN ¹³ CO	$(1621515 - 1521414)$	351548.3	316.1	6.17×10^{-4}	31
HN ¹³ CO	$(1621516 - 1521415)$	351548.3	316.1	6.17×10^{-4}	33
HN ¹³ CO	$(1621417 - 1521316)$	351561.8	316.1	6.19×10^{-4}	35
HN ¹³ CO	$(1621415 - 1521314)$	351561.8	316.1	6.17×10^{-4}	31
HN ¹³ CO				6.17×10^{-4}	33
	$(1621416 - 1521315)$	351561.8	316.1		

Table A.1. continued.

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Species	Transition	Frequency	$E_{\rm up}$	A_{ii}	$g_{\rm up}$
		(MHz)	(K)	(s^{-1})	
HN ¹³ CO	$(16\ 2\ 14\ 17 - 15\ 2\ 13\ 16)$	351561.8	316.1	6.19×10^{-4}	35
HN ¹³ CO	$(1621415 - 1521314)$	351561.8	316.1	6.17×10^{-4}	31
HN ¹³ CO	$(16\ 2\ 14\ 16 - 15\ 2\ 13\ 15)$	351561.8	316.1	6.17×10^{-4}	33
HN ¹³ CO	$(1601617 - 1501516)$	351642.9	143.5	6.30×10^{-4}	35
HN ¹³ CO	$(1601616 - 1501515)$	351642.9	143.5	6.27×10^{-4}	33
HN ¹³ CO	$(1601615 - 1501514)$	351642.9	143.5	6.27×10^{-4}	31

Table A.1. continued.

Notes. ^(a) This list only includes optically thin and unblended lines.

Appendix B: Additional figures

Fig. B.1. *Black*: Detected lines of NH₂CDO. *Red*: Best-fit model for T_{ex} =300 K.

Fig. B.2. *Black*: Detected lines of cis-NHDCHO. *Red*: Best-fit model for *T*ex=300 K.

Fig. B.3. *Black*: Detected lines of trans-NHDCHO. *Red*: Best-fit model for *T*ex=300 K.

Fig. B.4. *Black*: Detected lines of NH₂¹³CHO. *Red*: Best-fit model for T_{ex} =300 K. *Green*: Best-fit model for T_{ex} =100 K.

Fig. B.5. *Black*: Detected lines of DNCO. *Red*: Best-fit model for *T*ex=300 K.

Fig. B.6. *Black*: Detected lines of HN¹³CO. *Red*: Best-fit model for $T_{ex}=300$ K. *Green*: Best-fit model for $T_{ex}=100$ K.

Fig. B.7. *Black*: Lines of NH₂CHO $v=0$ with the lowest opacities. *Red*: Model based on the analysis of the NH₂¹³CHO lines and a ¹²C/¹³C ratio equal to 68.

Fig. B.8. *Black*: Line of NH₂CHO v_{12} =1 with the lowest opacity. *Red*: Model based on the analysis of the NH₂¹³CHO lines and a ¹²C/¹³C ratio equal to 68.