The physical and chemical structure of high-mass star-forming regions

Unraveling chemical complexity with the NOEMA large program "CORE"

C. Gieser^{1,*}, H. Beuther¹, D. Semenov^{1,2}, A. Ahmadi³, S. Suri¹, T. Möller⁴, M.T. Beltrán⁵, P. Klaassen⁶, Q. Zhang⁷, J.S. Urquhart⁸, Th. Henning¹, S. Feng^{9, 10, 11}, R. Galván-Madrid¹², V. de Souza Magalhães¹³, L. Moscadelli⁵, S. Longmore¹⁴, S. Leurini¹⁵, R. Kuiper¹⁶, T. Peters¹⁷, K.M. Menten¹⁸, T. Csengeri¹⁹, G. Fuller²⁰, F. Wyrowski¹⁸,

S. Lumsden²¹, Á. Sánchez-Monge⁴, L. Maud²², H. Linz¹, A. Palau¹², P. Schilke⁴, J. Pety^{13, 23}, R. Pudritz²⁴,

J.M. Winters¹³, and V. Piétu¹³

S. Longmore -, S. Leurin -, K. Kuiper -, I. Peiers -, K.M. Meinten -, I. Csenger -, C. Pull^{17, 2}, A. Padau 2; P. Schilke⁴, J. Pety^{13, 23}, R. Pudritz²⁴, J.M. Winters¹³, and V. Pićul¹³
 (Affiliations can be found after the references) Received...; accepted...
 ABSTRACT
 Amst. Characterizing the physical and chemical properties of forming massive stars at the spatial resolution of individual high-mass cores lies at the heart of current star formation research.
 Methods. We use sub-arcsecond resolution (-0.74) observations with the NOrthern Extended Millimeter Array at 1.37 mm to study the dust down to 300–2300 at that are resolved by our observations. We combine the derived physical and chemission. The density profiles are inferred from the 1.37 mm continuum visibilities. The column densities of 11 different species are determined by fitting the colusion and with a refacel by our observations. We combine the derived physical and chemical by fitting the colusion and with a resolute of 2000 at 1000 at that are resolved by our observations. We combine the derived physical and chemical by fitting the column densities of 11 different species are determined by fitting the colusion and with a radially decreas-indice of *q* = 0.4 ± 0.1 and an average density power-law index of *p* = 0.2 ± 0.2 using a point access on the 1.37 mm continuum wisibilities. The column densities of 11 different species are determined by fitting the correlations. We apply the physical-chemical model MUII stage Chemical. codE (MUSCLE) to the derived column densities of the cone and find a mean chemical age of -60000yrs and an age spread of 20000 - 1000 outyrs. With this paper we release all data products of the CORE project available at https://www.mpia.de/core.
 Conclusions. The CORE sample reveals well constrained density and temperature power-law distributions. Furthermore, we characterize a large warity in molecular richness that can be explained by an age spr

2007; Smith et al. 2009; Tan et al. 2014; Krumholz 2015; Schilke 2015; Motte et al. 2018; Rosen et al. 2020). High-mass stars are less common, located typically at large distances of several kiloparsecs and therefore difficult to observe at a high spatial resolution. The evolution of HMSF is fast, on the order of 10^5 yrs as revealed by observations (see, e.g., Table 2 in Motte et al. 2018; Mottram et al. 2011a) and theoretical models (McKee & Tan

frared and radio wavelengths. Various classifications exist in the literature, e.g., based on infrared properties, Cooper (2013) classifies massive young stellar objects (MYSOs) into three types: type I show strong H₂, but no ionized lines; type II are weaker in H₂, but show H_I emission in the Brackett series; type III have strong HI and weak H2 line emission. These near-infrared observations show that type III MYSOs are bluer than type I MYSOs indicating that they are more evolved.

In this study we follow the observationally driven nomenclature of four different evolutionary stages during HMSF by Gerner et al. (2014, 2015). When regions of a molecular cloud

^{*} Fellow of the International Max Planck Research School for Astronomy and Cosmic Physics at the University of Heidelberg (IMPRS-HD).

undergo gravitational collapse and fragmentation, protostars form in cold and dense clumps, often detected as infrared dark clouds (IRDCs, e.g., Rathborne et al. 2006). These clouds have typical temperatures of $\sim 10 - 20$ K and are visible at (sub)mm wavelengths (e.g., Pillai et al. 2006), but show no or only weak emission in the infrared. As the collapse continues, due to the conservation of angular momentum, a disk-outflow system forms around the protostar and the temperature of the surrounding envelope increases due to central heating by the protostar (e.g., Sánchez-Monge et al. 2013a; Beltrán & de Wit 2016). Through accretion of the infalling gas, protostars increase their masses and luminosities and become visible at infrared wavelengths. Protostars during this stage are referred to as high-mass protostellar objects (HMPOs, e.g., Sridharan et al. 2002; Williams et al. 2004; Beuther et al. 2010; Duarte-Cabral et al. 2013). HMPOs are characterized by high bolometric luminosities, $L > 10^4 L_{\odot}$, a strong thermal dust continuum, but only weak cm emission (Sridharan et al. 2002; Beuther et al. 2002). As the temperature of the surrounding envelope reaches $\gtrsim 100$ K, molecules that formed on the surfaces of dust grains and resided in ice layers, evaporate into the gas-phase revealing a rich molecular reservoir (e.g., Osorio et al. 1999). In this stage, the objects are referred to as hot molecular cores (HMCs) or "hot cores" (e.g., Cesaroni et al. 1997). HMPOs and HMCs have similar physical properties, however, the chemical composition of the gas around HMCs is richer due to higher temperatures in the envelope. Low-mass analogues are referred to as "hot corinos" (e.g., IRAS 16293-2422, Bottinelli et al. 2004). As the mass growth continues, the strong protostellar radiation dissociates and ionizes the surrounding envelope gas and an hyper/ultra-compact HII (HC/UCHII) region forms (e.g., Wood & Churchwell 1989; Hatchell et al. 1998; Churchwell 2002; Sánchez-Monge et al. 2013b). HII regions can be studied and classified through their strong free-free emission at cm wavelengths (e.g., Peters et al. 2010). These four evolutionary stages are not sharp transitions and overlap, especially between the intermediate HMPO/HMC stages and there are HMCs which have already formed a HC/UCHII region.

Aside from the physical complexity of HMSFRs, observations reveal one of the most complex chemical compositions in the interstellar medium (ISM, for a review, see, e.g., Herbst & van Dishoeck 2009; Jørgensen et al. 2020). To date about 200 molecules have been detected in the ISM (an overview is found in McGuire 2018), most of these can be observed toward the high-mass star-forming Galactic center molecular cloud Sagittarius B2 (e.g., Belloche et al. 2013). Spectroscopic studies at mm and sub-mm wavelengths have revealed the composition of the molecular gas in HMSFRs to be extremely diverse: nitrogen (N)-bearing species (such as HCN, CH₃CN); oxygen (O)-bearing species (such as H₂CO); sulfur (S)-bearing species (such as SO, SO₂). In shocked regions, enhanced abundances of S-bearing molecules such as SO₂ and silicon (Si)-bearing molecules such as SiO are observed (Schilke et al. 1997). During the HMC stage, a large variety of so-called complex organic molecules (COMs) evaporate into the gas-phase and produce line-rich spectra at mm wavelengths. We use the definition by Herbst & van Dishoeck (2009) in which a COM contains six or more atoms. In the densest regions where protostars form, the chemical composition of the molecular gas has been studied thoroughly. An important finding is the chemical segregation of O- and N- bearing species (e.g., Rodgers & Charnley 2001). This has been observed and studied toward many bright HMCs such as Orion-KL (Caselli et al. 1993; Feng et al. 2015); W3 H2O and W3 OH (Wyrowski et al. 1999; Qin et al. 2015); AFGL 2591

(Jiménez-Serra et al. 2012; Gieser et al. 2019); NGC7538 IRS9, W3 IRS5 and AFGL 490 (Fayolle et al. 2015); G35.20-0.74N (Allen et al. 2017); SgrB2(N) (Bonfand et al. 2017; Mills et al. 2018); or AFGL 4176 (Johnston et al. 2020).

In this paper, we want to pin down the physical and chemical properties of HMSFRs from dust and molecular line emission and with this information investigate the chemical timescales. We analyze the physical structure and molecular content of 18 HMSFRs in combination with a physical-chemical model on spatial scales ranging from 10 000 au down to our resolution limit of 300 au. We successfully tested the method presented in this paper on the well-studied hot core AFGL 2591 VLA3 (Gieser et al. 2019). In this case-study, we used the molecular line and dust continuum data from the observations of the CORE project to derive the density and temperature structure and molecular column densities of this hot core and applied a physical-chemical model to estimate the chemical age. Here, we apply this method to all 18 CORE regions with the goal to compare and understand the chemical diversity we observe during HMSF.

The paper is organized as followed: In Sect. 2 we summarize the observations, data calibration and imaging techniques. In Sect. 3 we derive the temperature and density structure of the regions. The molecular content is analyzed in Sect. 4. We apply a physical-chemical model to the observed molecular column densities in Sect. 5. The results of the physical and chemical properties of the regions are discussed in Sect. 6. A summary and our conclusions are given in Sect. 7.

2. Observations

The NOrthern Extended Millimeter Array (NOEMA) large program "Fragmentation and disk formation during high-mass star formation - CORE" is a high angular resolution survey (\sim 0."4 at 1.37 mm) designed to study the fragmentation, kinematic, and chemical properties of a homogeneous sample of 18 HMSFRs. The observations consist of spectral line and continuum data in Band 3 (1 mm). An overview of the project and results of the analysis of the dust continuum and fragmentation properties are presented in Beuther et al. (2018).

In combination with this paper, we release all data products of the CORE project with and without self-calibration conducted with GILDAS. The data can be found on the CORE collaboration website: https://www.mpia.de/core. The CLEANed data products as well as the calibrated *uv*-tables are available.

2.1. Sample

The sample of the CORE project consists of well-studied HMS-FRs on the Northern hemisphere. They were selected based on their bolometric luminosity ($L > 10^4 L_{\odot}$) and distance (d < 6 kpc) and allow us to study the evolution at early stages of HMSF in the HMPO/HMC stage. An overview of the properties of the regions is shown in Table 1 in this paper and additional properties, such as the luminosity L and mass M, are listed in Table 1 in Beuther et al. (2018). With declinations higher than +20°, most of the regions are difficult or impossible to observe with the Atacama Large Millimeter/submillimeter Array (ALMA), and therefore can only be studied with NOEMA at a high angular resolution (< 1") at mm wavelengths. The angular resolution for all regions is homogeneous (~0."4), but the resolved linear spatial scales vary from 300 au to 2 300 au, as the regions are located at distances between 0.7 kpc and 5.5 kpc.

Table 1: Overview of the CORE sample. The isotopic ratios are taken from Wilson & Rood (1994), further explained in Sect. 3.2, Sect. 4.2, and Sect. 5.1.

Region	Coord	linates	Distance	Galactic Distance	Velocity	Iso	otopic Rati	ios
	α	δ	d	$d_{ m gal}$	$v_{\rm LSR}$	$^{12}C/^{13}C$	$^{32}S/^{34}S$	¹⁶ O/ ¹⁸ O
	J(2000)	J(2000)	(kpc)	(kpc)	$({\rm km}~{\rm s}^{-1})$			
IRAS 23033	23:05:25.00	+60:08:15.5	4.3	10.4	-53.1	86	22	649
IRAS 23151	23:17:21.01	+59:28:47.5	3.3	9.8	-54.4	81	22	613
IRAS 23385	23:40:54.40	+61:10:28.0	4.9	11.1	-50.2	91	22	690
AFGL 2591	20:29:24.86	+40:11:19.4	3.3	8.2	- 5.5	69	22	519
CepA HW2	22:56:17.98	+62:01:49.5	0.7	8.4	-10.0	71	22	531
G084.9505	20:55:32.47	+44:06:10.1	5.5	9.4	-34.6	78	22	590
G094.6028	21:39:58.25	+50:14:20.9	4.0	9.3	+29.0	77	22	584
G100.38	22:16:10.35	+52:21:34.7	3.5	9.4	-37.6	78	22	590
G108.75	22:58:47.25	+58:45:01.6	4.3	10.3	-51.5	85	22	643
G138.2957	03:01:31.32	+60:29:13.2	2.9	10.5	-37.5	86	22	654
G139.9091	03:07:24.52	+58:30:48.3	3.2	10.8	-40.5	89	22	672
G075.78	20:21:44.03	+37:26:37.7	3.8	8.1	- 8.0	68	22	513
IRAS 21078	21:09:21.64	+52:22:37.5	1.5	8.3	- 6.1	70	22	525
NGC7538 IRS9	23:14:01.68	+61:27:19.1	2.7	9.5	-57.0	79	22	596
S106	20:27:26.77	+37:22:47.7	1.3	7.9	- 1.0	67	22	502
S87 IRS1	19:46:20.14	+24:35:29.0	2.2	7.3	+22.0	62	22	466
W3 H2O	02:27:04.60	+61:52:24.7	2.0	9.6	-48.5	80	22	602
W3 IRS4	02:25:31.22	+62:06:21.0	2.0	9.6	-42.8	80	22	602

Table 2: Overview of the CORE data products. The noise of the continuum data σ_{cont} is computed in a 3."5 × 3."5 rectangle in an emission-free region. I_{peak} is the continuum peak intensity. The average map noise of the merged (combined NOEMA + IRAM 30m) spectral line data $\sigma_{line,map}$ is computed in a line-free range from a rest-frequency of 219.00 GHz to 219.13 GHz with a channel width of 3 km s⁻¹ (2.2 MHz) within the full primary beam.

Region	Sy	/nthesi	ized Beam	Synthesized Beam			CASA		GILDAS		
	Continuum	Data	Merged Line	e Data	Standar	d Calibration	Self-calibration		Self-calibra		ation
	$\theta_{\rm maj} \times \theta_{\rm min}$	PA	$\theta_{\rm maj} \times \theta_{\rm min}$	PA	$\sigma_{ m cont}$	Ipeak	$\sigma_{ m cont}$	Ipeak	$\sigma_{ m cont}$	Ipeak	$\sigma_{ m line,map}$
	("×")	$(^{\circ})$	(''×'')	(°)	(mJ	y beam ⁻¹)	(mJy	beam ⁻¹)	(mJy	beam ⁻¹)	(K)
IRAS 23033	0.45×0.37	47	0.43×0.43	96	0.50	28.21	0.43	38.76	0.53	38.13	0.40
IRAS 23151	0.45×0.37	50	0.47×0.38	54	0.26	26.41	0.26	32.65	0.25	33.78	0.48
IRAS 23385	0.48×0.43	58	0.49×0.44	55	0.29	14.61	0.16	18.03	0.16	17.84	0.30
AFGL 2591	0.47×0.36	65	0.48×0.37	65	0.53	56.31	0.50	87.33	0.51	85.10	0.65
CepA HW2	0.42×0.41	41	0.44×0.39	80	3.72	239.93	2.39	441.07	2.43	448.50	0.68
G084.9505	0.43×0.38	69	0.44×0.39	72	0.10	4.37	0.08	6.28	0.08	5.19	0.28
G094.6028	0.41×0.39	77	0.41×0.38	85	0.15	10.24	0.15	13.65	0.13	12.84	0.36
G100.38	0.49×0.33	56	0.49×0.34	55	0.07	6.35	0.05	8.51	0.06	7.28	0.25
G108.75	0.50×0.44	49	0.51×0.44	49	0.17	10.53	0.12	14.75	0.13	16.08	0.27
G138.2957	0.50×0.41	60	0.51×0.41	59	0.16	6.28	0.16	6.15	0.10	6.28	0.33
G139.9091	0.51×0.40	56	0.52×0.41	56	0.18	9.23	0.20	13.84	0.15	13.49	0.34
G075.78	0.48×0.37	60	0.49×0.37	60	0.47	46.31	0.38	64.50	0.38	71.67	0.58
IRAS 21078	0.48×0.33	41	0.48×0.33	41	0.41	26.76	0.44	34.64	0.44	36.69	0.41
NGC7538 IRS9	0.44×0.38	80	0.44×0.38	80	0.32	29.14	0.19	41.28	0.21	46.09	0.42
S106	0.47×0.34	47	0.48×0.34	47	0.88	77.41	0.96	136.00	1.20	150.47	0.35
S87 IRS1	0.54×0.35	37	0.55×0.36	37	0.12	19.42	0.17	33.35	0.18	37.37	0.44
W3 H2O	0.43×0.32	86	0.45×0.32	86	4.28	279.11	2.54	449.84	2.82	456.29	0.74
W3 IRS4	0.45×0.32	82	0.46×0.32	83	0.69	21.73	0.42	39.34	0.33	51.50	0.72

The regions show a large diversity of fragmentation properties (Beuther et al. 2018): while some regions contain mainly a single isolated core (e.g., AFGL 2591), other regions fragment into up to 20 cores (e.g., IRAS 21078). Individual case-studies were carried out on some of the regions, e.g., the kinematic properties of W3 H2O (Ahmadi et al. 2018), IRAS 23385 (Cesaroni et al. 2019), IRAS 23033 (Bosco et al. 2019), W3 IRS4 (Mottram et al. 2020), and IRAS 21078 (Moscadelli et al. 2021); the chemical composition of the pilot regions NGC7538 S and NGC7538 IRS1 (Feng et al. 2016), and AFGL 2591 (Gieser et al. 2019). A multi-wavelength modeling study of AFGL 2591 is presented in Olguin et al. (2020). In Ahmadi et al. (in prep.) the kinematic analysis of the complete CORE sample will be covered, while in this study we focus on the physical structure (temperature and density) and chemical composition of the molecular gas. We do not include the pilot regions NGC7538 S and NGC7538 IRS1 presented in Beuther et al. (2018) in our analysis, as only for the remaining 18 regions we have a homogeneous



Fig. 1: Comparison of the GILDAS standard calibrated (left panel), CASA self-calibrated (middle panel) and GILDAS self-calibrated (right panel) W3 IRS4 continuum data. In each panel, the beam size is shown in the bottom left corner and the pink bar in the top left corner indicates a linear spatial scale of 5 000 au.



Fig. 2: *Top panels:* Comparison of the GILDAS standard calibration (left) and GILDAS self-calibration (right) of the W3 IRS4 broad-band spectral line data zoomed in toward the position of the continuum peak. The integrated intensity of the CH₃CN $12_3 - 11_3$ transition is shown in color. In each panel, the beam size is shown in the bottom left corner and pink bar in the top left corner indicates a linear spatial scale of 5 000 au. *Bottom panel:* CH₃CN 12_3-11_3 spectrum at the position of the W3 IRS4 continuum peak of the GILDAS standard calibrated data (black) and of the GILDAS self-calibrated data (green). The dashed orange line shows the systemic velocity of the region v_{LSR} and the dash-dotted blue lines show the lower and upper velocity limits ($v_{LSR} \pm 10 \text{ km s}^{-1}$) used for the integrated intensity map shown in the top panel.

multi-configuration NOEMA data set. The pilot studies have no corresponding D array observations, so less *uv*-coverage, and could hence not be accurately compared. However, a detailed individual chemical analysis of the pilot regions is already presented in Feng et al. (2016).

2.2. NOEMA Observations

The sample was observed from 2014 to 2017 in the A (extended), the old B or new C (intermediate), and D (compact) array configurations. Two regions each were observed in track-sharing pairs to reduce calibration time. Spectral line data were obtained with the broad-band WideX correlator at a rest-frequency range of 217.2 – 220.8 GHz with a spectral resolution of ~2.7 km s⁻¹. In addition, eight narrow-band units were placed within this frequency range in order to obtain high spectral resolution data (~0.4 km s⁻¹) of kinematically interesting lines such as CH₃CN (a summary of the high spectral resolution units is summarized in Table 2 and 3 in Ahmadi et al. 2018). The interferometric data were calibrated with the CLIC package in GILDAS¹. The 1.37 mm continuum data were extracted from the broad-band WideX data by carefully selecting line-free channels in each region.

2.3. IRAM 30m Observations

Interferometers spatially filter the extended emission. The shortest baseline of the NOEMA array is $\sim 20 \text{ m}$, thus spatial scales larger than 16" are not recovered by the interferometric observations at 1.37 mm. All CORE regions were therefore observed with the IRAM 30m telescope using the Eight MIxer Receiver (EMIR, Carter et al. 2012) in order to recover the missing flux in the spectral line data due to spatial filtering. EMIR has four basebands and each have a width of ~4 GHz and a spectral resolution of 200 kHz corresponding to ~ 0.3 km s⁻¹ at 1.37 mm). The lower inner baseband (LI) of our EMIR spectral setup covers the same spectral range as the NOEMA observations with the broad-band WideX correlator. The half power beam width (HPBW) of the IRAM 30m telescope is 11."8 at 1.37 mm. For a detailed description of the IRAM 30m data calibration we refer to Appendix A in Mottram et al. (2020). The continuum data have no complementary single-dish observations, so here spa-

¹ https://www.iram.fr/IRAMFR/GILDAS/



Fig. 3: 1.37 mm continuum emission. The selected positions analyzed in this paper (summarized in Table A.1) are marked and labeled in red (core positions) and blue (all remaining positions). The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with contour steps increasing by a factor of 2 (e.g., -5, 5, 10, 20, 40, 80,... σ_{cont} , see Table 2 for values of σ_{cont} for each region). In each panel, the beam size is shown in the bottom left corner and the pink bar in the top left corner indicates a linear spatial scale of 5 000 au. The field of view in each region is adjusted to only show the area with significant emission ($\geq 5\sigma_{cont}$), and for regions with wide-spread emission, the primary beam is indicated by a black circle.

tial filtering can remain an issue as discussed in Beuther et al. 2.4. Sel (2018).

2.4. Self-calibration

The majority of the continuum data have a high signal-to-noise (S/N) ratio in the GILDAS standard calibrated data (see Table 2). However, the phase noise can be high due to an unstable atmosphere during the observations causing the flux to be scat-



Fig. 3 (cont.): 1.37 mm continuum emission. The selected positions analyzed in this paper (summarized in Table A.1) are marked and labeled in red (core positions) and blue (all remaining positions). The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with contour steps increasing by a factor of 2 (e.g., -5, 5, 10, 20, 40, 80,... σ_{cont} , see Table 2 for values of σ_{cont} for each region). In each panel, the beam size is shown in the bottom left corner and the pink bar in the top left corner indicates a linear spatial scale of 5 000 au. The field of view in each region is adjusted to only show the area with significant emission ($\geq 5\sigma_{cont}$), and for regions with wide-spread emission, the primary beam is indicated by a black circle.

tered around the source. This issue can be improved by applying phase self-calibration to the interferometric data (e.g., Pearson & Readhead 1984; Radcliffe et al. 2016).

The CORE continuum data presented in Beuther et al. (2018) were successfully phase self-calibrated using the Common Astronomy Software Applications package (CASA) through an iterative masking of the source. However, at that time it was not possible to apply the self-calibration solution to the spectral line

data. The CASA phase self-calibration of the continuum data was performed by applying an interactive mask in each self-calibration loop starting with the strongest structures first and proceeding with the weaker structures (Beuther et al. 2018). Depending on the S/N ratio of the region, solution intervals of 220, 100, or 45 s were used.

We have now used the self-calibration tool in GILDAS to phase self-calibrate the continuum as well as the spectral line



Fig. 3 (cont.): 1.37 mm continuum emission. The selected positions analyzed in this paper (summarized in Table A.1) are marked and labeled in red (core positions) and blue (all remaining positions). The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with contour steps increasing by a factor of 2 (e.g., -5, 5, 10, 20, 40, 80,... σ_{cont} , see Table 2 for values of σ_{cont} for each region). In each panel, the beam size is shown in the bottom left corner and the pink bar in the top left corner indicates a linear spatial scale of 5 000 au. The field of view in each region is adjusted to only show the area with significant emission ($\geq 5\sigma_{cont}$), and for regions with wide-spread emission, the primary beam is indicated by a black circle.

data of the CORE project to provide homogeneously calibrated data products of the full CORE observations. The crucial point of phase self-calibration is to start with a good enough spatial model of the source. The selfcal procedure of the GILDAS/MAPPING package uses as a source model the first CLEAN components n_{CLEAN} found during a previous step of deconvolution. The basic idea is that the first CLEAN components deliver a model of the source with a high S/N ratio whose spatial structure is not much affected by flux scattering. The number of CLEAN components n_{CLEAN} must be large enough to get a fair



Fig. 3 (cont.): 1.37 mm continuum emission. The selected positions analyzed in this paper (summarized in Table A.1) are marked and labeled in red (core positions) and blue (all remaining positions). The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with contour steps increasing by a factor of 2 (e.g., -5, 5, 10, 20, 40, 80,... σ_{cont} , see Table 2 for values of σ_{cont} for each region). In each panel, the beam size is shown in the bottom left corner and the pink bar in the top left corner indicates a linear spatial scale of 5 000 au. The field of view in each region is adjusted to only show the area with significant emission ($\geq 5\sigma_{cont}$), and for regions with wide-spread emission, the primary beam is indicated by a black circle.

representation of the source structure and small enough to avoid deconvolving scattered flux that would be confused with noise. The visibilities are usually averaged in time to increase their signal-to-noise ratio during the first self-calibration iteration and this averaging time is progressively lowered to the minimum possible integration time in the following iterations. In practice, we started to deconvolve the continuum source with an absolute flux stopping criterion set to 3 times the continuum noise. This gives a number of CLEAN iterations n_{iter} that we used to iterate the self-calibration three times increasing n_{CLEAN} from $n_{\text{iter}}/4$, to



Fig. 3 (cont.): 1.37 mm continuum emission. The selected positions analyzed in this paper (summarized in Table A.1) are marked and labeled in red (core positions) and blue (all remaining positions). The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with contour steps increasing by a factor of 2 (e.g., -5, 5, 10, 20, 40, 80,... σ_{cont} , see Table 2 for values of σ_{cont} for each region). In each panel, the beam size is shown in the bottom left corner and the pink bar in the top left corner indicates a linear spatial scale of 5 000 au. The field of view in each region is adjusted to only show the area with significant emission ($\geq 5\sigma_{cont}$), and for regions with wide-spread emission, the primary beam is indicated by a black circle.

 $n_{\text{iter}}/2$, and to n_{iter} , and decreasing the averaging time from 200 s, to 100 s, and to 45 s. Only visibilities with a S/N ratio > 6 are self-calibrated, but the remaining visibilities are kept in the proceeding CLEANing of the data in order to not loose visibilities on the longest baselines which would decrease the angular resolution. With this method, simple structures, as well as regions with a complicated morphology can be successfully phase self-calibrated. As the continuum data has the highest sensitivity, we used the solution from the continuum self-calibration and apply the gain solution to the broad-band and narrow-band spectral line data using the UV_CAL task.

An example on how self-calibration is increasing the quality of the continuum data quality is shown in Fig. 1 for W3 IRS4 which has a complex morphology in the continuum data. While GILDAS standard calibrated data already reveals the complex structure of the region, the noise is high throughout the primary beam with many negative features. Applying the CASA and GILDAS self-calibration significantly lowers the noise and increases the peak intensity. The continuum noise and peak intensity are summarized for all regions in Table 2. The mean continuum noise $\sigma_{\rm cont}$ is 0.74 mJy $\rm beam^{-1}$ in the standard calibrated data, while for the CASA and GILDAS self-calibrated data, the mean noise $\sigma_{\rm cont}$ is lowered by ~25% to 0.53 mJy beam⁻¹ and 0.56 mJy beam⁻¹, respectively. In general, $\sigma_{\rm cont}$ is higher toward the strong continuum sources S106, CepA HW2, and W3 H2O. The mean S/N ratio of the GILDAS standard calibrated continuum data is 74, while for the CASA and GILDAS self-calibrated continuum data, the mean S/N is improved by a factor of two to 130 and 132, respectively.

Even though both CASA and GILDAS self-calibration methods improve the data quality, due to the different techniques to define a source model (interactive masking and defining number of clean components, respectively), there are differences in the resulting images. In Fig. 1, a faint structure toward the NW of the continuum peak is seen in the GILDAS standard calibrated and self-calibrated image, but it is not recovered in the CASA selfcalibrated image. Comparing the self-calibration results (Table 2), the peak intensities are higher in the GILDAS self-calibrated data, while a lower noise is achieved in the CASA self-calibrated data. For very faint structures and a complex source morphology, a careful interpretation of the self-calibrated data is recommended, but overall all the main features are recovered with both self-calibration methods.

A comparison between the standard and self-calibrated broad-band spectral line data is shown in Fig. 2 for the CH_3CN $12_3 - 11_3$ transition around the location of the 1.37 mm continuum peak in the W3 IRS4 region. The emission is less fuzzy and more compact in the self-calibrated data product. On individual spectra, self-calibration provides a significant increase of the line intensity which has a big impact, e.g., when deriving column densities.

2.5. Data products and public release

The data merging of the interferometric and single-dish data and imaging is done in the GILDAS/MAPPING package. The WideX and the corresponding EMIR spectral-line data are smoothed to a common spectral resolution of 3.0 km s^{-1} . The narrow-band and the corresponding EMIR spectral line data are smoothed to a common spectral resolution of 0.5 km s^{-1} . In order to merge the NOEMA and IRAM 30m data, the task UVSHORT converts the short spacings into a pseudo *uv*-table and combines them with the NOEMA data. The deconvolution of the NOEMA continuum, NOEMA spectral line and merged (combined NOEMA + IRAM 30m) spectral line data is performed in GILDAS/MAPPING using the Clark CLEAN algorithm (Clark 1980) adopting three different weightings: robust weighting with a robust parameter of 0.1 ($\theta \sim 0.''4$), a robust parameter of 1.0 ($\theta \sim 0.''6$), and natural weighting ($\theta \sim 1.''0$). The stopping criterion for the continuum and broad-band spectral line data is set to $f_{\rm res} = 0.01$ which corresponds to a minimum fraction of 1% of the peak intensity in the dirty image. The stopping criterion for the narrow-band spectral line data is either $n_{\rm iter} = 5\,000$ (maximum number of iterations) or $a_{\rm res} = 0.01$, which corresponds to a maximum intensity of 10 mJy beam⁻¹ in the residual image.

Molecules such as CO isotopologues and SO may have extended emission in the merged data products (Mottram et al. 2020). As the Clark algorithm assumes emission from point sources, the CLEANed map may have point-like artifacts. The SDI algorithm (Steer et al. 1984) may improve the CLEANed image in such cases. A comparison between the Clark and SDI algorithms for imaging of the W3 IRS4 region is presented in Mottram et al. (2020). Final data products of the merged broadband spectral line data of molecular lines with potential large-scale emission (13 CO 2 – 1, SO 6₅ – 5₄, H₂CO 3_{0,3} – 2_{0,2}, H₂CO 3_{2,2} – 2_{2,1}, H₂CO 3_{2,1} – 2_{2,0}) CLEANed using the SDI algorithm with robust weighting (robust parameter of 3, $\theta \sim 0."6$) with a stopping criterion of $n_{iter} = 25\,000$ are provided as well. Primary beam correction is applied to the continuum and spectral line data products.

In this study we use the primary beam corrected NOEMAonly continuum and merged (NOEMA + IRAM 30m) broadband spectral line data. Both data products are imaged with the Clark algorithm and a robust parameter of 0.1 resulting in the highest angular resolution ($\theta \sim 0.''4$). Table 2 summarizes the properties of the standard and self-calibrated data products for all regions including the synthesized beam (major axis θ_{maj} , minor axis θ_{min} , and position angle PA), noise of the continuum data σ_{cont} , continuum peak intensity I_{peak} , and noise in the merged (NOEMA + IRAM 30m) spectral line data $\sigma_{line,map}$. The mean map noise of the merged spectral line data is 0.44 K.

3. Physical structure

The linear spatial resolution of the CORE sample ranges from 300-2300 au. At this spatial resolution it is not possible to resolve potential disks surrounding the protostars but these can be studied and characterized through the kinematic analysis of the line profiles (Ahmadi et al. 2019). However, we do resolve the gas and dust envelopes around individual cores which can be approximated as spherically symmetric objects for which the radial temperature profile T(r) and radial density profile n(r) of the envelope gas can be described by power-laws (e.g., van der Tak et al. 2000; Beuther et al. 2002; Palau et al. 2014):

$$T(r) = T_{\rm in} \times \left(\frac{r}{r_{\rm in}}\right)^{-q} \tag{1}$$

$$n(r) = n_{\rm in} \times \left(\frac{r}{r_{\rm in}}\right)^{-p} \tag{2}$$

with $T_{in} = T(r_{in})$ and $n_{in} = n(r_{in})$ at an arbitrary radius r_{in} . The temperature power-law index q and density power-law index p of the core envelopes are important properties of HMSFRs. By studying these quantities with observations, theoretical analytical models on how massive stars form can be constrained.

Article number, page 10 of 81

3.1. Continuum emission

A detailed analysis of the continuum data is presented in Beuther et al. (2018). The updated GILDAS self-calibrated continuum data are shown in contours in Fig. 3. While the sample was selected to be largely homogeneous in luminosity, at an angular resolution of ~0."4 a variety of structures can be identified. While some regions appear as isolated single objects (IRAS 23151, AFGL 2591), others show spatially separated cores (IRAS 23033, G108.75, G139.9091, S106, CepA HW2, W3 H2O). There are regions in which fragmentation is observed within an embedded envelope (IRAS 23385, IRAS 21078). Many of the regions have a complex morphology such as filamentary structures and extended envelopes (G084.9505, G094.6028, G100.38, G138.2957, G75.78, NGC7538 IRS9, S87 IRS1, W3 IRS4).

In this study we aim to analyze the physical properties and chemical variation across the regions, and have therefore selected a number of positions throughout the different regions. Deriving the column density of all detected species in every single spectrum in each region is computationally expensive and we restrict this method to the analysis of the H₂CO and CH₃CN temperature maps (Sect. 3.2). In order to study the cores and envelopes around forming protostars we select positions which have a clear spherically symmetric core-like morphology in the 1.37 mm continuum data which is the case for most regions toward the continuum peak position, but also multiple spherically symmetric objects within a region can be identified (e.g., toward IRAS 23033). In addition, we select positions in the extended envelopes to study potential differences in the chemical abundances. The in total 120 selected positions are summarized in Table A.1 and marked in Fig. 3. The nomenclature of each position is denoted by the region name and increasing number with decreasing 1.37 mm continuum intensity. For each region, position 1 is toward the 1.37 mm continuum emission peak.

In order to estimate the H_2 column density (Sect. 4.1), we require that all positions are detected in the 1.37 mm continuum $(I_{1.37\text{mm}} \ge 5\sigma_{\text{cont}})$. The number of selected positions is higher toward regions with extended envelopes and complex morphologies (e.g., IRAS 21078 and G138.2957) compared to compact regions (e.g., G139.9091 and S106). In this study, we define in Sect. 3.2 a "core" as an object that shows a radially decreasing temperature profile over at least the width of two beams. In Sect. 5, we will apply a 1D physical-chemical model to these defined cores using the observed temperature and density structure analyzed in Sect. 3.2 and Sect. 3.3, respectively and observed molecular column densities (Sect. 4). These positions are labeled as "C" (core) in Table A.1. The remaining positions are locations in the dust envelope and environment around the cores. There are a few positions with a clear spherically symmetric core morphology in the dust emission, but no temperature profile can be derived (e.g., position 1 in S106). Cores with unresolved radial temperature profiles are also not included in this approach. In total we select 120 positions including 22 core positions. The broad-band spectra of the 120 positions are shown in Fig. F.1. Table A.1 lists the noise of the broad-band spectrum extracted from these positions and the systemic velocity v_{LSR} determined from the $C^{18}O 2 - 1$ line, which may differ from the average region v_{LSR} listed in Table 1 due to velocity gradients within the region (see Sect. 4.2). In contrast to this core definition, in the analysis by Beuther et al. (2018), "cores" are defined as fragmented objects with emission $\geq 10\sigma_{cont}$ using the clumpfind algorithm (Williams et al. 1994) and thus more cores are found in their analysis.



Fig. 4: Radial temperature profiles of the 22 cores. Each panel shows the binned radial temperature profile derived from the H₂CO (green) and CH₃CN (blue) temperature maps shown in Fig. C.1. The data points used for the radial profile fit are shown by corresponding colored errorbars, the data points excluded from the fit are indicated by grey errorbars. The outer radius of the temperature fit is shown by the vertical purple dash-dotted line. The inner unresolved region is shown as a grey-shaded area. The linear fit and the $\pm 1\sigma$ uncertainty are shown by the solid and dashed red lines, respectively.

3.2. Temperature structure

To reliably determine the rotation temperature T_{rot} , it is required to observe multiple transitions of a molecule. We use formalde-

hyde (H_2CO) and methyl cyanide (CH_3CN) as thermometers to determine the temperature structure of the regions, as both have multiple strong and optically thin lines in our spectral setup. The spectral line properties of these molecules are listed in Table B.1.



Fig. 4 (cont.): Radial temperature profiles of the 22 cores. Each panel shows the binned radial temperature profile derived from the H_2CO (green) and CH_3CN (blue) temperature maps shown in Fig. C.1. The data points used for the radial profile fit are shown by corresponding colored errorbars, the data points excluded from the fit are indicated by grey errorbars. The outer radius of the temperature fit is shown by the vertical purple dash-dotted line. The inner unresolved region is shown as a grey-shaded area. The linear fit and the $\pm 1\sigma$ uncertainty are shown by the solid and dashed red lines, respectively.

We model the spectral line emission of these molecules using the eXtended CASA Line Analysis Software Suite (XCLASS,

Möller et al. $2017)^2$. With XCLASS, molecular lines can be modeled and fitted by solving the 1D radiative transfer equation. In

² https://xclass.astro.uni-koeln.de/

Table 3: Results of the physical structure (Sect. 3) and estimated chemical ages (Sect. 5) of the cores. The density power-law index p is calculated according to Eq. (3). The core mass M_{core} is calculated according to Eq. (5). The estimate of the chemical timescale τ_{chem} is explained in detail in Sect. 5.2.

Region +	r _{in}	r _{out}	$T_{\rm kin}(r_{\rm in})$	\overline{q}	α	p	M _{core}	$ au_{ m chem}$
Number	(au)	(au)	(K)				(M_{\odot})	(yrs)
IRAS 23033 1	1 837	5 7 2 0	114.9 ± 8.2	$0.34 \pm 0.11^{\dagger}$	-0.44 ± 0.03	2.22 ± 0.11	6.06±1.29	3.4(4)-9.8(4)
IRAS 23033 2	1837	4767	167.2 ± 6.6	$0.48 {\pm} 0.20^{\dagger}$	-0.73 ± 0.05	1.79 ± 0.21	7.81±1.59	1.8(4)
IRAS 23033 3	1837	3813	160.8 ± 8.0	$0.96 {\pm} 0.07^{\dagger}$	-0.65 ± 0.05	1.39 ± 0.09	5.22 ± 1.08	1.9(4)
IRAS 23151 1	1 392	2926	129.2 ± 4.7	$0.27 \pm 0.11^*$	-0.48 ± 0.01	2.25 ± 0.11	3.28 ± 0.67	1.9(4) - 8.4(4)
IRAS 23385 1	2 299	4 3 4 5	239.5±12.1	$0.14 \pm 0.05^{\dagger}$	-0.63 ± 0.05	2.23 ± 0.07	4.43 ± 0.92	4.9(4)
IRAS 23385 2	2 299	4 3 4 5	226.0 ± 2.3	$0.21 \pm 0.04^{\dagger}$	-0.39 ± 0.06	2.40 ± 0.07	2.30 ± 0.46	9.5(4)
AFGL 2591 1	1 394	2926	159.9 ± 6.8	$0.25 \pm 0.08^*$	-0.63 ± 0.02	2.12 ± 0.08	7.43±1.52	8.4(4)
CepA HW2 1	289	931	238.2±11.4	$0.31 \pm 0.02^*$	-0.44 ± 0.02	2.25 ± 0.03	1.24 ± 0.26	8.4(4)
CepA HW2 2	289	776	170.5 ± 13.3	$0.56 \pm 0.10^{*}$	-0.45 ± 0.05	1.99 ± 0.11	0.28 ± 0.06	1.9(4) - 8.8(4)
G084.9505 1	2273	6 0 97	169.0 ± 0.8	$1.38 \pm 0.05^{\dagger}$	-0.79 ± 0.03	0.83 ± 0.06	1.67 ± 0.33	1.7(4)
G094.6028 1	1 584	4434	183.8 ± 45.4	$1.48 \pm 0.27^{\dagger}$	-0.75 ± 0.04	0.77 ± 0.27	2.35 ± 0.76	5.7(4) - 8.4(4)
G100.38 1	1 4 5 2	3 880	68.2 ± 0.7	$0.41 \pm 0.22^{\dagger}$	-0.76 ± 0.05	1.83 ± 0.23	1.85 ± 0.37	8.4(4)
G108.75 1	2044	4767	111.2 ± 8.2	$0.56 \pm 0.09^{\dagger}$	-0.30 ± 0.03	2.14 ± 0.09	2.58 ± 0.55	2.0(4)
G108.75 2	2044	5720	82.9 ± 2.6	$0.23 \pm 0.02^{\dagger}$	-0.87 ± 0.04	1.90 ± 0.04	3.73 ± 0.76	1.1(5)
G075.78 1	1642	5 0 5 5	176.8 ± 0.3	$0.29 \pm 0.05^*$	-0.64 ± 0.04	2.07 ± 0.06	9.21±1.84	8.4(4)
IRAS 21078 1	612	1 3 3 0	135.5 ± 0.2	$0.43 \pm 0.05^*$	-0.90 ± 0.05	1.67 ± 0.07	1.60 ± 0.32	5.5(4) - 9.2(4)
IRAS 21078 2	612	1 3 3 0	121.9 ± 3.4	$0.59 \pm 0.01^*$	-1.10 ± 0.05	1.31 ± 0.05	1.70 ± 0.34	4.9(4)
NGC7538 IRS9 1	1114	2 3 9 4	200.5 ± 0.9	$0.38 \pm 0.04^*$	-0.37 ± 0.04	2.25 ± 0.06	1.60 ± 0.32	8.4(4)
S87 IRS1 1	1 0 0 5	2439	112.0 ± 5.2	$0.45 \pm 0.07^{\dagger}$	-0.52 ± 0.04	2.03 ± 0.08	2.15 ± 0.44	5.9(4)
W3 H2O 3	770	1774	176.6±10.9	$0.59 \pm 0.06^{*}$	-0.65 ± 0.09	1.76 ± 0.11	11.61 ± 2.44	8.4(4)
W3 H2O 4	770	1774	166.4 ± 9.0	$0.45 \pm 0.03^*$	-0.89 ± 0.09	1.66 ± 0.09	11.22 ± 2.33	8.6(4)
W3 IRS4 1	780	1 774	173.0±15.4	$0.54 \pm 0.08^{*}$	-0.52 ± 0.05	1.94 ± 0.09	1.14 ± 0.25	2.0(4) - 8.6(4)

Notes. $a(b) = a \times 10^b$. Fit results for r_{in} , r_{out} , $T_{kin}(r_{in})$, and q derived from either the H₂CO or CH₃CN radial temperature profiles (Fig. 4) are indicated by either a \dagger or *, respectively.



Fig. 5: Histogram of the temperature power-law index q. The red line shows a Gaussian fit to the data points for q = 0.1 - 0.6.

the calculation of the Gaussian line profiles optical depth effects are included. For each molecule, the properties of all transitions are taken from an embedded SQlite3 database containing entries from the Cologne Database for Molecular Spectroscopy (CDMS, Müller et al. 2005) and the Jet Propulsion Laboratory (JPL, Pickett et al. 1998) using the Virtual Atomic and Molecular Data Centre (VAMDC, Endres et al. 2016).

The broad-band spectral setup includes three strong formaldehyde lines, two of which have the same upper energy level, and one weak transition from the $H_2^{13}CO$ isotopologue. H_2CO and $H_2^{13}CO$ are fitted simultaneously in XCLASS using an

isotopic ratio calculated from Wilson & Rood (1994): ${}^{12}C/{}^{13}C \approx$ 7.5 × d_{gal} + 7.6. The ${}^{12}C/{}^{13}C$ ratio is listed in Table 1 for each region. While formaldehyde is a good low-temperature gas tracer at $T_{kin} < 100$ K (Mangum & Wootten 1993), at high densities and temperatures the $3_{0,3} - 2_{0,2}$ transition is optically thick and a reliable temperature cannot be derived anymore from the line ratios with this method (Rodón et al. 2012; Gieser et al. 2019). To determine temperatures at higher densities, we use nine methyl cyanide lines (J = 12 - 11 and K = 0 - 8, $E_u/k_B = 69 - 526$ K). In addition, 7 lines of the CH₁₃¹³CN isotopologue are fitted simultaneously (J = 12 - 11 and K = 0 - 6, $E_u/k_B = 69 - 326$ K). A detailed discussion of the line optical depth is given in Sect. 4.2.

In XCLASS, each molecule can be described by a number of emission and absorption components. The fit parameter set of each component consists of the source size θ_{source} , the rotation temperature T_{rot} , the column density N, the linewidth Δv , and the velocity offset from the systemic velocity v_{off} . One can choose a variety of algorithms that can also be combined in an algorithm chain in order to find the best-fit parameters by minimizing the χ^2 value. We adopt an algorithm chain with the Genetic and Levenberg-Marquardt (LM) methods optimizing toward global and local minima, respectively.

For each region, we use the myXCLASSMapFit function to fit the H₂CO and CH₃CN lines in each pixel within the primary beam. All lines with a peak intensity > $10\sigma_{\text{line,map}}$ are fitted with one emission component. We chose a high threshold of $10\sigma_{\text{line,map}}$ so multiple transitions have a high signal-to-noise ratio to accurately determine the rotation temperature. Only a single value can be set as the threshold in the myXCLASSMapFit function, therefore toward the edges of the primary beam the



Fig. 6: Radial visibility profiles of the 1.37 mm continuum emission of the 22 cores. The black data points show the radial profile of the averaged complex visibilities of the 1.37 mm continuum as a function of uv distance (bottom x-axis) and of the corresponding linear scale (top x-axis). The linear fit and the $\pm 1\sigma$ uncertainties are indicated by the solid and dashed red line, respectively.

Article number, page 14 of 81



Fig. 6 (cont.): Radial visibility profiles of the 1.37 mm continuum emission of the 22 cores. The black data points show the radial profile of the averaged complex visibilities of the 1.37 mm continuum as a function of uv distance (bottom x-axis) and of the corresponding linear scale (top x-axis). The linear fit and the $\pm 1\sigma$ uncertainties are indicated by the solid and dashed red line, respectively.



Fig. 7: Histogram of the density power-law index p. The density power-law index derived with the measured values of q for each core are shown in blue. In green, the results for p calculated with a fixed temperature index (q = 0.4) for all cores are shown. The red line shows a Gaussian fit to the green histogram.

temperature estimates are less reliable due to an increase of the noise.

Under the assumption of local thermal equilibrium (LTE), the kinetic temperature of the gas can be estimated from the rotation temperature $T_{\rm kin} \approx T_{\rm rot}$. As HMSFRs have high densities $n > 10^5 \,{\rm cm}^{-3}$ toward the locations of the protostars, LTE can be assumed here (Mangum & Shirley 2015). The critical densities $n_{\rm crit} = \frac{A_{\rm ul}}{C_{\rm ul}}$ for the CH₃CN and H₂CO lines are ~4×10⁶ cm⁻³ and ~3×10⁶ cm⁻³, respectively ($A_{\rm ul}$ is the Einstein A coefficient and $C_{\rm ul}$ is the collisional rate coefficient). Here, we use $C_{\rm ul}$ measured at 140 K taken from the Leiden Atomic and Molecular Database (LAMDA, Schöier et al. 2005). The H₂CO and CH₃CN temperature maps of each region are shown in Fig. C.1. The H₂CO lines trace the extended low temperature gas at 10 – 100 K. The CH₃CN maps mostly show spatially compact emission tracing the higher temperature gas at >100 K.

For each selected position (Table A.1), we extract the H₂CO and CH₃CN radial temperature profile which are binned in steps of half a beam (Δr). In each bin, the uncertainties are computed from the standard deviation of the mean. After a first visual inspection of all radial profiles of the 120 positions, we select all positions with a radially decreasing temperature profile along at least two beams in at least one temperature tracer. With this method, we are able to extract in total 22 positions, which we define as "cores". The radial temperature profiles of the cores are shown in Fig. 4. In cases where both temperature tracers are detected toward the central core (e.g., IRAS 23033 2), the observed H₂CO temperature profile has significantly higher values compared to CH₃CN. As discussed previously, this is due to the fact that due to the high optical depth in these dense regions, the fit algorithm converges toward high temperatures. The line optical depth is discussed further in Sect. 4.2.

We fit the profiles with a power-law according to Eq. (1) using the minimum χ^2 method to derive the temperature powerlaw index q. We use the CH₃CN temperature profile for the fit if it is detected along at least two beams and the H₂CO temperature profile otherwise. The inner radius is the temperature at a radius of half the beam size $r_{\rm in} = \Delta r$. The outer radius $r_{\rm out}$ is chosen as a local minimum, when $T_{\rm rot}(r_{\rm out}) < T_{\rm rot}(r_{\rm out} + \Delta r)$ and $T_{\rm rot}(r_{\rm out}) < T_{\rm rot}(r_{\rm out} + 2\Delta r)$. In case the outer radii of nearby cores would overlap or the 2D temperature distribution becomes

Article number, page 16 of 81

highly asymmetric, we reduce the outer radius. The outer radii are also marked in Fig. 4. The fit results for q, r_{in} , r_{out} , and $T_{kin}(r_{in})$ are summarized in Table 3 for each core.

A histogram of the temperature power-law index q is shown in Fig. 5. Most of the data points are distributed between 0.1 and 0.6, but three outliers are located at q > 0.9. A Gaussian fit to the data with q = 0.1 - 0.6 yields an average value of $q = 0.4 \pm 0.1$, which is in very good agreement with theoretical predictions (Emerson 1988; Osorio et al. 1999; van der Tak et al. 2000) and observations (e.g., Palau et al. 2014, 2020). The issue of optical depth of the H₂CO lines and CH₃CN being only detected toward the densest parts result in the high uncertainties and spread in q. We observe a broad range of q with shallow profiles (q = 0.1) up to steep profiles (q = 1.5). In order to study if q is constant for all high-mass cores or if the range of q is real, observations of more cores are required. Osorio et al. (1999) and van der Tak et al. (2000) suggest steeper values for q on scales $< 2\,000$ au. In addition, the temperature maps and radial profiles are 2D projected, and the real 3D profile may be more complicated.

3.3. Density structure

In contrast to the merged (NOEMA + IRAM 30m) spectral line data, the 1.37 mm continuum NOEMA data suffer from missing flux due to the lack of short-spacing information. Therefore, it is difficult to reliably derive radial intensity profiles in the final images that are required to determine the density structure. This issue can be minimized by analyzing instead the complex visibilities of the 1.37 mm continuum data in the *uv*-plane (Adams 1991; Looney et al. 2003; Beuther et al. 2007b; Zhang et al. 2009). Assuming spherical symmetry, the power-law index of the complex visibility profile α is related to the density powerlaw index of the radial density distribution *p* by (Looney et al. 2003):

$$\alpha = p + q - 3. \tag{3}$$

For each core position, the phase center is shifted to this location. The remaining cores within a region are fitted as a point source + circular Gaussian (to account both for the compact and the extended emission) and subtracted using the UV_FIT task in GILDAS. With this approach blending of nearby cores in the visibility profiles within a single region can be minimized. The azimuthally averaged complex visibilities computed using the UVAMP task in MIRIAD (Sault et al. 1995) are binned in steps of $20 k\lambda$. The radial visibility profiles are shown in Fig. 6. The uv distances range from $\sim 20 - 800$ m corresponding to spatial scales of $\sim 10^3 - 10^5$ au at distances of a few kpc. We apply a power-law fit to the data in order to derive the power-law index α . The density power-law index p is calculated according to Eq. (3) taking into account the temperature power-law index q derived in Sect. 3.2. The results for α and p for all cores are summarized in Table 3.

Most of the visibility profiles (Fig. 6) can be described by a single power-law. The higher scatter at large uv distances is due to the fact that less data is available at long baselines. For IRAS 23385, the visibility profiles of core 1 and 2 do not follow a simple power-law. Toward smaller spatial scales (< 10⁴ au), the profiles are steeper. This could be due to the fact, that even though the contribution of nearby cores is minimized by subtracting a point source + circular Gaussian, in the case of IRAS 23385 it is not possible to clearly distinguish the contribution of both cores, which are embedded within a common dust envelope. A histogram of the density index p is shown in Fig. 7. Using the observationally derived values of q, the power-law index ranges from 1.4 – 2.4. Fixing the temperature power-law index q to the mean value of 0.4 and calculating p with the results from the *uv*-analysis, the distribution of p gets narrower, peaking at p = 2.0 - 2.1. A Gaussian fit to this distribution yields a mean of $p = 2.0 \pm 0.2$. The results of the derived physical structure of these 22 cores are discussed in detail in Sect. 6.1.

4. Molecular gas content

In this section, we analyze the chemical content of the molecular gas toward the 18 CORE HMSFRs by studying the molecular column densities derived toward the 120 positions. The column density of molecular hydrogen $N(H_2)$ is derived from the 1.37 mm dust continuum emission (Sect. 4.1). In addition, molecular column densities are derived from the merged spectral line data with XCLASS by fitting the emission lines assuming LTE conditions (Sect 4.2). Spectra extracted toward the selected positions are shown in Fig. F.1. In total, we consider 11 species among in total 16 isotopologues commonly detected within the 4 GHz spectral bandwidth of the broad-band WideX correlator: ¹³CO, C¹⁸O, SO, OCS, SO₂, DCN, H₂CO, HNCO, HC₃N, CH₃OH, CH₃CN. A spectral resolution of 3 km s⁻¹ is not sufficient to study the line widths and kinematic properties in detail, but sufficient to derive molecular column densities *N*.

4.1. Molecular hydrogen and core mass estimate

The beam-convolved molecular hydrogen column density $N(H_2)$ toward all 120 positions and mass calculation of the 22 cores $M_{\rm core}$ can be derived from the continuum intensity I_{ν} assuming that the emission is optically thin (Hildebrand 1983):

$$N(\mathrm{H}_2) = \frac{I_{\nu}\eta}{\mu m_{\mathrm{H}}\Omega\kappa_{\nu}B_{\nu}(T)},\tag{4}$$

$$M_{\rm core} = \frac{F_{\nu} \eta d^2}{\kappa_{\nu} B_{\nu}(T)},\tag{5}$$

with a gas-to-dust mass ratio $\eta = 150 \ (\eta = \frac{M_{\text{gas}}}{M_{\text{H}}} / \frac{M_{\text{dust}}}{M_{\text{H}}}$, with $\frac{M_{\text{gas}}}{M_{\text{H}}} = 1.4$ and $\frac{M_{\text{dust}}}{M_{\text{H}}} = 0.0091$, see Table 1.4 and 23.1 in Draine 2011, respectively), mean molecular weight $\mu = 2.8$ (Kauffmann et al. 2008), hydrogen mass m_{H} , beam solid angle Ω , dust opacity $\kappa_{\nu} = 0.9 \text{ cm}^2 \text{ g}^{-1}$ for dust grains with a thin icy mantle at a gas density of 10^6 cm^{-3} at 1.3 mm (Ossenkopf & Henning 1994), the Planck function $B_{\nu}(T)$ and distance *d*. F_{ν} is the integrated intensity of each core within the outer radius r_{out} .

We use $T_{\rm kin}$ for the temperature T in the Planck function taken from the thermometers H₂CO and CH₃CN, assuming LTE conditions ($T_{\rm kin} \approx T_{\rm rot}$). If the spectrum is extracted from a core position, $T_{\rm kin}$ is taken from the radial temperature fit described in Sect. 3.2 with $T = T_{\rm kin}(r_{\rm in})$ (see Table 3). If the spectrum is extracted from a position not corresponding to a core, the kinetic temperature is computed from $T_{\rm rot}$ (CH₃CN) if detected or from $T_{\rm rot}$ (H₂CO) otherwise. If there is no temperature tracer detected toward the position, we adopt a lower limit of $T_{\rm kin} = 20 \pm 10$ K, as the lowest derived rotation temperatures range between 10–30 K.

In order to validate that the assumption of optically thin dust emission is valid, the continuum optical depth τ_{ν}^{cont} is computed

for each position (for a derivation of the equation, see Appendix A in Frau et al. 2010):

$$\tau_{\nu}^{\text{cont}} = -\ln\left(1 - \frac{I_{\nu}}{\Omega B_{\nu}(T)}\right). \tag{6}$$

The kinetic temperature T_{kin} , molecular hydrogen column density $N(H_2)$, and continuum optical depth τ_{ν}^{cont} are listed in Table A.1 for all 120 positions. The uncertainties of $N(H_2)$ and $M_{\rm core}$ are calculated assuming Gaussian error propagation and include the uncertainty of the continuum intensity with an estimated 20% flux calibration uncertainty (Beuther et al. 2018) and uncertainty of the derived rotation temperature ΔT_{kin} (listed in Table A.1). The optical depth τ_{ν}^{cont} is $\ll 1$ toward most positions, so optically thin dust emission can be assumed here and the H₂ column density and core mass calculation provide reliable results. The only exceptions with $\tau_{\nu}^{\text{cont}} > 1$ are the positions 1 and 2 of the W3 H2O region which corresponds to the W3 OH UCHII region (see also Ahmadi et al. 2018). The molecular hydrogen column density $N(H_2)$ has a mean value of 1.5×10^{24} cm⁻², ranging from 2.7×10^{22} cm⁻² up to 2.8×10^{25} cm⁻². The core masses $M_{\rm core}$ are listed in Table 3. We find a mean core mass of 4.1 M_{\odot} in a range between $0.3 M_{\odot}$ and $11.6 M_{\odot}$. As discussed previously, due to missing short-spacing information, both $N(H_2)$ and M_{core} should be considered as lower limits. However, the high core masses indicate that they are harboring protostars which will eventually end up as massive stars.

4.2. Spectral line modeling with XCLASS

We use the spectral line data of the CORE project to derive molecular column densities of 11 different species using the XCLASS software. A description of the XCLASS software is presented in Sect. 3.2. Using the myXCLASSFit function, individual spectra are fitted species by species with one emission component to derive the molecular column density N.

A spectrum is extracted from the merged spectral line data for each position listed in Table A.1. The noise σ_{line} in each spectrum is computed in a line-free range from 219.00 GHz to 219.13 GHz. Compared to the average map noise $\sigma_{\text{line,map}}$ listed in Table 1, the noise in each spectrum σ_{line} may have higher or lower noise values since the noise distribution is not uniform within the primary beam. The systemic velocity v_{LSR} is determined by fitting the C¹⁸O 2 – 1 transition and may differ from the global v_{LSR} listed in Table 1 due to velocity gradients in the region, hence it allows us to employ a narrow parameter range for v_{off} so fitting strong nearby emission lines is avoided. The systemic velocity v_{LSR} and noise σ_{line} are listed in Table A.1 for each position.

All molecules and their corresponding transitions that are fitted with XCLASS are listed in Table B.1. Lines which are blended with transitions of other detected species (at a resolution of 3 km s^{-1}) are also listed, but excluded from the fit. For most molecules, only the rotational ground-state level v = 0can be detected in our spectral setup. For SO₂ and HC₃N vibrationally excited levels ($v_x > 0$) are present and for CH₃OH torsionally excited lines are detected ($v_t = 1$). The following species for which we observe multiple isotopologues are fitted simultaneously: OCS and O¹³CS, SO₂ and ³⁴SO₂, H₂CO and H¹³₂CO, HC₃N and HCC¹³CN, CH₃CN and CH¹³₃CN. The isotopologue ratios are summarized in Table 1 and are calculated either from Wilson & Rood (1994) ($^{12}C/^{13}C \approx 7.5 \times d_{gal} + 7.6$) or taken from references within ($^{32}S/^{34}S \approx 22$). The uncertainties of these isotopic ratios are high due to a large scatter of the



Fig. 8: Column density histograms. The H_2 column density is derived from the 1.37 mm continuum emission and the remaining molecular column densities are derived with XCLASS. Column density histograms of all 120 positions are shown in green bars (upper limits are not included). Separate column density histograms of the core and the remaining positions are indicated by the dash-dotted red and dashed blue lines, respectively.

data points. However, we do not observe a sufficient number of strong isotopic lines to measure it more precisely. We do not fit 13 CO and C 18 O simultaneously, because the 13 CO 2 – 1 line has a high optical depth (see Table B.1).

We use an algorithm chain with the Genetic and Levenberg-Marquardt (LM) methods and to estimate the uncertainties of the fit parameters, we use the MCMC error estimation algorithm afterward. The following criteria are applied to the fitted model spectrum and column density $N_{-\Delta N_{low}}^{+\Delta N_{upp}}$ of each species in order to determine bad fits: 1) model spectra with a peak intensity $< 3\sigma_{\text{line}}$; 2) the upper error of the column density ΔN_{upp} is converging towards high values ($\Delta N_{upp} > 10 \times N$); and 3) the lower error of the column density is not constrained $\Delta N_{low} = 0 \text{ cm}^{-2}$. With these three criteria, weak and unresolved lines are automatically discarded and we use the best-fit value of *N* only as an upper limit.

The column densities N for all species fitted with XCLASS and their uncertainties derived with the MCMC error estimation algorithm are summarized in Tables E.1 - E.3 for all positions. Histograms of the logarithmic column density distributions including $N(H_2)$ are shown in Fig. 8 with the mean and standard variation of the column density noted in each panel (upper limits are not included). The logarithmic bin width is set to 0.5. Separate histograms of the core and non core populations are shown in the same panels. However, as discussed in Sect. 2, for the spectral line data we have short-spacing information, while we do not have this for the continuum data, hence we systematically underestimate the H₂ column density.

There are species with a distribution having a clear column density peak (e.g., H_2 , ¹³CO, C¹⁸O, SO, DCN, H_2 CO, HNCO). Other species have a double-peaked distribution with a clear separation between the core and the remaining positions (e.g., OCS, HC₃N, CH₃OH, CH₃OH; $v_t = 1$, and CH₃CN). In these cases, the column density is enhanced by a factor of ~ 10 – 100 toward the core positions. There are not enough data points for the SO₂, HC₃N; $v_7 = 1$, HC₃N; $v_7 = 2$ to draw any conclusions about the distribution, however, they are detected mostly in the densest regions toward core positions. In general, high column densities are found toward the remaining positions.

To account for the fact that toward the core positions the column density is expected to be higher in a higher density region, we show relative abundance $N(X)/N(C^{18}O)$ histograms in Fig. D.1 (upper limits are not included). Assuming that both species trace the same emission region, the relative abundances are independent of the absolute column density value, which differ from region to region and from core to core. Normalized to $N(C^{18}O)$, most species have a single-peaked distribution. Exceptions are OCS, HNCO, HC₃N, CH₃OH, and CH₃CN with double-peaked distributions indicating that high temperature gas-phase chemistry has a big impact on these N-bearing molecules by increasing their abundances. However, in general there is still a clear difference between core positions (high abundance) and non core positions (low abundance). The fact that larger molecules have a clearer distinction between the core and non core positions, while for simpler species it is less obvious (e.g., ¹³CO and DCN), hints that the emission of COMs is associated with the cores while simple molecules are abundant in envelope as well. The difference of the core and the remaining positions indicates that the high densities and possible energetic processes around the protostars have a strong impact on the molecular abundances in the gas-phase (e.g. through protostellar outflows, shocks, disk accretion, and strong radiation from the protostars). Correlations of the derived column densities are discussed in Sect. 6.3.

Observed and XCLASS modeled spectra are shown in Fig. F.1 for all positions. The computed optical depth for all fitted lines as a function of rest-frequency is shown as well. Even though

the sample was selected to be at a similar evolutionary stage (HMPO/HMC), the number of emission lines in the observed spectra vary from region to region, but also within a region. Typical hot core spectra are observed for the positions AFGL 2591 1, CepA HW2 1, G75.78 1, W3 H2O 1, and W3 H2O 2. Many weak emission features are detected in the spectra at a ~ $2-3\sigma_{\text{line}}$ level originating from COMs. These COM emission features are difficult to fit as the transitions are weak, have similar upper energy levels $E_{\rm u}/k_{\rm B}$, and are blended at a spectral resolution of 3 km s⁻ so they were excluded from this analysis. A detailed study of the integrated line emission (including line stacking of weak COM emission lines) will be presented in a future study (Gieser et al. in prep.). Fewer species are detected in spectra toward the non core positions. In contrast to the line-rich sources, there are several sources that show only a handful of emission lines mainly from CO-isotopologues, SO, and H₂CO even toward the continuum peak positions. Line-poor regions are G139.9091, G138.2957, S87 IRS1, and S106. In the case of IRAS 23033, core 1 has a line-poor spectrum, while core 2 and 3, which are embedded in a common envelope, significantly more emission lines are detected. These cores either could be at different evolutionary stages or are embedded in an inhomogeneous local radiation field. The former can be investigated by applying a chemical model to the observed column densities and by estimating the chemical ages of the regions, which we investigate in the next section.

The mean and maximum line optical depth, $\tau_{\text{mean}}^{\text{line}}$ and $\tau_{\text{max}}^{\text{line}}$, computed for each fitted transition with XCLASS are summarized in Table B.1. The ¹³CO 2 – 1 transition has the highest mean optical depth with $\tau_{\text{mean}}^{\text{line}} = 1.5$, but for most other species and transitions, the mean line optical depth is < 1, so the column density and temperature determination should be reliable, especially when many transitions of a molecule are fitted simultaneously. The mean optical depth of the H₂CO 3_{0,3} – 2_{0,2} transition is a factor of 4 higher than the remaining two transitions. When the 3_{0,3} – 2_{0,2} transition is optically thick, temperature estimates depending on the line ratios are difficult to determine with the remaining two optically thin lines as they have similar upper energy levels (68 K). In 23 out of the 118 spectra where H₂CO was detected and fitted with XCLASS, the calculated optical depth of the 3_{0,3} – 2_{0,2} transition is $\tau_{\text{line}} > 1$.

5. Physical-chemical modeling of the cores

The continuum data of the CORE sample show a large diversity in fragmentation properties (Beuther et al. 2018) and our analysis in Sect. 4 showed that the composition of the molecular gas varies within as well as between the regions: some have a rich plethora of molecular lines, while others have line-poor spectra. This diversity of physical and chemical properties could be explained by a number of reasons. Magnetic fields and/or different initial density structures could explain the variety in fragmentation properties (Beuther et al. 2018). Different initial conditions in, e.g., the large-scale kinematics and mass distribution might also have an effect on the molecular abundances. To investigate if the observed variation of the physical and chemical properties of the cores may be due to the fact that the CORE regions have different ages, we model the chemical evolution of the 22 cores in the following.

5.1. MUSCLE setup

A physical-chemical model is applied to the physical properties and molecular column densities of each core determined from



Fig. 9: Comparison of the observed and modeled column densities shown in red and blue, respectively. Upper limits are indicated by an arrow.

the CORE 1 mm observations in order to estimate the chemical ages. MUSCLE (MUlti Stage ChemicaL codE) was already successfully applied to the CORE pilot regions NGC7538S and NGC7538IRS1 (Feng et al. 2016) and the CORE region



Fig. 9 (cont.): Comparison of the observed and modeled column densities shown in red and blue, respectively. Upper limits are indicated by an arrow.

AFGL 2591 (Gieser et al. 2019). The model comprises spherically symmetric physical structures. The temperature and den-

sity profiles of the cores are described by power-laws up to the outer radius r_{out} with index q and p, respectively, see Eq. (1) and

Fabl	le 4:	MUSC	CLE	input	para	ameters.
------	-------	------	-----	-------	------	----------

Parameter	Value
Radiation field:	value
CP ionization rate ζ_{rec}	$1.8(-16)s^{-1}a$
Extinction A at r	1.0(-10)s 10^{mag}
LIV photodesorption yield	$1(-05)^{b,c}$
Crain properties:	1(-03)
Grain radius r	$0.1 \mu m^d$
Dust density α	$0.1 \mu \text{m}$
Dust defisity ρ_d	150 ^e
Casto-dust mass ratio η	130
Surface diffusivity $E_{\text{Diff}}/E_{\text{Bind}}$	0.4 ³
Mantle composition	olivine
Initial Chemical Abundances	hast ft IDDC stage
HNPO model	12 700 stage °
Inner radius $r_{\rm in}$	12 /00 au
Outer radius r_{out}	0.5 pc
Temperature $T_{\rm in}$ at $r_{\rm in}$	11.3 K
Temperature power-law index q	0.0 (isothermal)
Density $n_{\rm in}$ at $r_{\rm in}$	$1.4(5) \mathrm{cm}^{-3}$
Density power-law index p	1.5
Stage lifetime $\tau_{\rm IRDC}$	16 500 years
HMC model	best-fit HMPO stage ^g
Inner radius $r_{\rm in}$	103 au
Outer radius r_{out}	0.5 pc
Temperature $T_{\rm in}$ at $r_{\rm in}$	75.8 K
Temperature power-law index q	0.4
Density $n_{\rm in}$ at $r_{\rm in}$	$1.5(9) \mathrm{cm}^{-3}$
Density power-law index p	1.8
Stage lifetime $ au_{ m HMPO}$	32 000 years
UCHII model	best-fit HMC stage ^g
Inner radius <i>r</i> _{in}	1 140 au
Outer radius r_{out}	0.5 pc
Temperature T_{in} at r_{in}	162.9 K
Temperature power-law index q	0.4
Density $n_{\rm in}$ at $r_{\rm in}$	$1.3(8) \mathrm{cm}^{-3}$
Density power-law index p	2.0
Stage lifetime $\tau_{\rm HMC}$	35 000 years

Notes. $a(b) = a \times 10^{b}$.

^(a) Indriolo et al. (2015); ^(b) Cruz-Diaz et al. (2016); ^(c) Bertin et al. (2016); ^(d) Gerin (2013); ^(e) Draine (2011); ^(f) Cuppen et al. (2017); ^(g) Table A.4, A.5, and A.6 in Gerner et al. (2015).

(2). At an inner radius r_{in} and further in, the density and temperature reach a constant value. We adopt 40 logarithmic grid points for the radial profiles.

On top of this static physical structure, the time-dependent gas-grain chemical network ALCHEMIC (Semenov et al. 2010) computes the abundances of hundreds of atomic and molecular species using thousands of reactions. A detailed description of MUSCLE can be found in Gerner et al. (2014, 2015). We adopt most of the model parameters from the AFGL 2591 case-study described in Gieser et al. (2019) which yielded a good estimate of the chemical age of this hot core compared to literature estimates. A summary of the input parameters is listed in Table 4. In contrast to the AFGL 2591 case-study, we use a higher value for the cosmic ionization rate ζ_{CR} based on a study of multiple HMSFRs by Indriolo et al. (2015). These authors find that ζ_{CR} is constant at a Galactic radius > 5 kpc and with all the CORE regions at Galactic distances > 7 kpc (Table 1), we use a constant value of $1.8 \times 10^{-16} \text{ s}^{-1}$. By setting the extinction at r_{out} to

 $A_v = 10^{mag}$, the core is shielded from the interstellar ultraviolet radiation field.

For each of the 22 cores, we run a physical-chemical model with MUSCLE. The input are the H₂ column density (Sect. 4.1) and all molecular column densities derived with XCLASS (Sect. 4.2). The CO column density N(CO) is calculated from $N(C^{18}O)$ as $C^{18}O$ is less optically thick than ¹³CO (Table B.1) and hence more reliably fitted in XCLASS. For each region, we calculate the ¹⁶O/¹⁸O isotopic ratio according to Wilson & Rood (1994): ¹⁶O/¹⁸O $\approx 58.8d_{gal} + 37.1$. The ¹⁶O/¹⁸O ratio is listed in Table 1 for each region. For HC₃N and CH₃OH we compute the mean column density of the rotational ground state and vibrationally/torsionally excited states for the MUSCLE input. Molecular column densities, for which only upper limits could be determined, are also set as upper limits in MUSCLE. We set the temperature structure of the model core to the observed temperature profile (Sect. 3.2) and use the density power-law index derived from the continuum visibility analysis (Sect. 3.3).

Two undetermined model parameters remain. First, we do not know how evolved the cores are, described by the chemical age $\tau_{\rm chem}$, and second, what the initial chemical composition of the parental molecular cloud/clump was. Due to the fact that the CORE regions are far more evolved than typical cold IRDCs and the physical structure of each model stage is static, one has to define a sensible initial condition for the chemical composition. The initial conditions we apply are based on a study of 69 HMS-FRs using single-dish observations (Gerner et al. 2014, 2015). These HMSFRs were classified according to their evolutionary stage (IRDCs, HMPOs, HMCs, and UCHII regions) and a template was created from the average column densities for each evolutionary stage. The four template stages were modeled using MUSCLE to create average abundances for all molecular and atomic species and to estimate a mean chemical age of each evolutionary stage. The properties of their template IRDC, HMPO, and HMC model are summarized in Table 4. Following the convention by Gerner et al. (2014, 2015), the chemical age τ_{chem} is 0 yrs when the gas density reaches 10^4 cm^{-3} .

Based on the temperature profiles of the 22 cores, we can assume that they lie somewhere between the HMPO and early UCHII stage, since the average temperatures around cores are too high to be classified as IRDCs ($T \sim 20$ K). There are a few known UCHII regions with strong free-free emission at cm wavelengths resolved in the CORE data. In W3 H2O, the Western part (around position 1 and 2 in Fig. 3) is the UCHII region W3 OH. In W3 IRS4, the Southern ring-like structure is an UCHII region as well (Mottram et al. 2020). However, for the W3 OH UCHII region we do not find a clear radial decreasing temperature profile and toward the W3 IRS4 UCHII region no H₂CO or CH₃CN line emission is detected at a $10\sigma_{\mathrm{line,map}}$ level to estimate the kinetic temperature. The S106 region is a UCHII as well for which we do not detect neither H₂CO nor CH₃CN emission around the compact core. This already suggests that toward this later stage the molecular richness in these regions is decreased.

To test which initial conditions (see Table 4) fit best to the observed molecular column densities, we model each of the 22 cores with initial abundances after an initial IRDC phase (referred to as the HMPO model), after an initial HMPO phase (referred to as the HMC model), and after an initial HMC phase (referred to as the UCHII model). While most cores are unlikely to have formed a strong UCHII region yet, we include the UCHII model, as the observations in Gerner et al. (2014, 2015) have large beam sizes (11" and 29"), and UCHII regions may have contamination from less evolved line-rich objects, which are blended into the single-pointing spectra. It is not our aim to

classify the cores into these evolutionary stages, but find sensible initial chemical conditions as an input for our physical-chemical modeling. For example, an evolved HMC, that is more evolved than the template HMC from Gerner et al. (2015), will be described best by the UCHII model in our nomenclature.

For each model, the chemical evolution τ_{model} runs up to 100 000 years in 100 logarithmic time steps. In each time step, the computed radial abundance profiles are converted into beam-convolved column densities with the beam size fixed to the mean synthesized beam of the observations. The best-fit model is determined by a minimum χ^2 analysis by comparing the modeled and observed column densities in each time step and for all three adopted initial condition models. Applying this physical-chemical model allows us to estimate the chemical age.

5.2. Chemical ages

The best-fit chemical age τ_{chem} , χ^2 , and percentage of well modeled molecules are shown in Table G.1 for each initial abundance model and core. Gerner et al. (2014) estimate that chemical ages are uncertain by a factor of 2 – 3 and that the modeled column densities are uncertain by a factor of 10. But this depends on the number of modeled molecules, but also cores embedded in complex dynamic environments are harder to fit with our model. The chemical age τ_{chem} is the sum of the time of the initial abundance model and τ_{model} , so for the HMPO model: $\tau_{chem} = \tau_{IRDC} + \tau_{model}$, for the HMC model: $\tau_{chem} = \tau_{IRDC} + \tau_{HMPO} + \tau_{model}$, and for the UCHII model: $\tau_{chem} = \tau_{IRDC} + \tau_{HMPO} + \tau_{model}$.

For some cores, multiple initial condition models have a similarly low χ^2 (e.g., the HMPO and UCHI model for core 1 in IRAS 23033). In these cases, we cannot constrain the chemical age well. Comparing the lowest χ^2 model with the remaining initial condition models, if the χ^2 difference is less than 5%, only chemical age ranges spanning over these models are further considered. Table 3 shows the chemical age τ_{chem} for models with a clear lowest χ^2 initial condition model or a time range in chemical age for cores with multiple best-fit initial condition models. The estimated chemical timescales τ_{chem} of the cores vary between ~20 000–100 000 yrs within the regions of the CORE sample with a mean of ~60 000 yrs. The youngest core being G084.9505 1 and the oldest core being G108.75 2.

A comparison of the best-fit modeled and observed column densities for all cores is shown in Fig. 9. For most cores, the model underestimates the H₂CO and CH₃OH column densities compared to the observed values. This can be explained by the fact that the quasi-static model does not sufficiently take into account the warm-up stage from 30 - 80 K where surface chemistry on the dust grains plays an important role and where these two molecules are formed by subsequent hydrogenation of CO. These discrepancies between modeled and observed H₂CO and CH₃OH column densities have already been noticed by Gerner et al. (2014) in their template HMPO stage modeled with MUSCLE. They explain that this is due to the fact that the formation route of H₂CO consists of grain-surface as well as gas-phase chemistry which are both time-dependent and not correctly implemented in the chemical models. This results in the over and underproduction of these species, which is also the case in our modeling results shown in Fig. 9.

Large discrepancies between the modeled and observed column density exist also for the SO molecule for which the model overproduces the SO column density by a factor > 10 for many cores (e.g., IRAS 23033 core 1, 2, and 3). This overproduction in SO is seen in all three initial condition models, but in most cases, other modeled S-bearing species (OCS and SO₂) are modeled well. The applied initial chemical conditions based on the Gerner et al. (2015) models also included S-bearing species (SO, CS, and OCS). Their initial IRDC stage model started with elemental abundances and only H₂ in molecular form taken from the low metals set of Lee et al. (1998). But in order to fit the IRDC phase accurately, Gerner et al. (2015) had to increase the initial elemental S abundance from 8×10^{-8} to 8×10^{-7} (w.r.t. H). However, an overproduction of SO is also seen in their best-fit HMPO, HMC and UCHII models. This might be connected to a poorly understood chemistry of the reactive SO molecule, as also in their models the remaining S-bearing species can be reproduced properly. In addition, as only one SO transition is covered in our spectral setup, which can be typically optically thick (Table B.1 and Fig. F.1), we may underestimate the observed SO column density. This might partially explain the differences between the modeled and observed SO column density.

With multiple cores resolved within a region, it is possible to study how the chemical timescale τ_{chem} varies across small spatial scales. In the IRAS 23033 region, core 1 seems to be more evolved ($\sim 30\,000 - 100\,000\,\mathrm{yrs}$), even though the spectrum is line-poor (see Fig. F.1) compared to the spectra of core 2 and 3 which are embedded in a common envelope (see Fig. 3) and for which we estimate similar chemical timescales of ~20 000 yrs. Core 1 and 2 in CepA HW2 have a chemical age of ~80 000 yrs and ~ $20\,000 - 90\,000$ yrs, respectively. The cores are very close (~2300 au), but within our sensitivity limit, these cores are not embedded in a common envelope, but have very steep density profiles ($p \ge 2$). In IRAS 21078, core 1 and 2 have a chemical age of $\sim 60\,000 - 90\,000$ yrs and $\sim 50\,000$ yrs, respectively, suggesting a small age gradient. The cores are embedded within a common envelope and have small projected separations. Core 3 and 4 in W3 H2O have a chemical age of ~90 000 yrs and $\sim 20\,000 - 90\,000$ yrs, respectively. In the IRAS 23385 region, core 1 is younger (~50 000 yrs), while core 2 is estimated to be much older (~100 000 yrs). In G108.75 a large difference between the chemical ages of core 1 (~20000 yrs) and core 2 (\sim 110000 yrs) is estimated. The cores have a separation of $\sim 20\,000$ au, but have the same systemic velocity (Table A.1). A strong external radiation field or complex dynamics could be the reason for this large chemical age difference.

One of the limitations of MUSCLE is that the physical structure (radial temperature and density profiles) is static within each evolutionary stage (IRDC, HMPO, HMC and UCHII). In reality these properties do change on timescales smaller than the chemical timescales derived here, and also the dynamics (e.g., gas inflow) are important factors to consider. Currently, 3D timedependent physical models in combination with a full chemical network are computationally expensive. Therefore, we use the approach of our quasi-static physical model by considering the four different evolutionary stages. More sophisticated physicalchemical models in the future are required to include 3D gas dynamics and the evolution of the density and temperature structure. In addition, a larger number of molecular column densities would better constrain the model parameter space.

6. Discussion

6.1. Physical structure of high-mass star-forming cores

Various methods were applied in the literature to observationally derive the density profiles of envelopes in HMSFRs (e.g., summarized in Table 6 in Gieser et al. 2019). Some of these studies are based on observations with single-dish telescopes with beam sizes > 10'' tracing the clump-scale envelope, while in-



Fig. 10: Literature comparison of the density index p at different core/clump sizes $\langle r \rangle$. Studies based on interferometric observations are marked by a "×", (sub)mm single-dish observations by a "•", multi-wavelength observations by a "•", and mid-infrared observations by a "o". Notes. M02: Mueller et al. (2002), B02: Beuther et al. (2002), H03: Hatchell & van der Tak (2003), W05: Williams et al. (2005), B07: Beuther et al. (2007b), Z09: Zhang et al. (2009), B12: Butler & Tan (2012), G13: Giannetti et al. (2013), P14: Palau et al. (2014), W16: Wyrowski et al. (2016), P20: Palau et al. (2020).



Fig. 11: Spearman correlation coefficient r for the derived physical and chemical core parameters listed in Table 3. Values higher than 0.8 are marked by a green circle.

terferometric observations trace the core-scale envelope. We select studies in the literature for which the density structure was determined for a sample of cores or clumps and also extract the typical sizes < r > from their studies. The results in comparison with our study are shown in Fig. 10. At scales of 1 pc down to 0.01 pc, it seems that the density index *p* lies between 1.7 and 2.0, which is close to the values inferred in low-mass star-forming regions (Motte & André 2001). To investigate this further, we have observed the CORE regions with the NIKA2 instrument at the IRAM 30m telescope and an analysis of the density structure at clump-scales will follow (Beuther et al. in prep.).

The observationally-derived density and temperature profiles $(q = 0.4 \pm 0.1 \text{ and } p = 2.0 \pm 0.2)$ are in agreement with theoretical studies of HMSF, but the physics of how massive stars form



Fig. 12: The M/L ratio and chemical ages τ_{chem} . The mass M and luminosity L for each region are taken from Beuther et al. (2018). The horizontal dashed lines correspond to average M/L ratios for HMPOs (red) and UCHII regions (blue) taken from Sridharan et al. (2002).

is not fully understood vet. Currently theoretical models propose the formation of high-mass stars through: a monolithic collapse of turbulent cores (McKee & Tan 2002, 2003); protostellar collisions and coalescence in dense clusters (Bonnell et al. 1998; Bonnell & Bate 2002); or competitive accretion in clusters (Bonnell et al. 2001; Smith et al. 2009; Hartmann et al. 2012; Murray & Chang 2012). The density and temperature structure are important parameters of the initial cloud and proceeding clump and core scales. For example, early star formation models by Shu (1977) and Shu et al. (1987) that model the gravitational collapse of an isothermal sphere find that p = 2 in the outer envelope and p = 1.5 in the inner region where the gas is free-falling onto the central region. McLaughlin & Pudritz (1996, 1997) used a logatropic equation of state and a non-isothermal sphere and find that at an initial density profile of p = 1, the profile steepens to p = 1.5 after the collapse. In the turbulent core model by McKee & Tan (2002, 2003) the authors assume p = 1.5 based on observational constraints. In Bonnell et al. (1998) the density profile in the outer region has the form p = 2 and a shallower, nearuniform profile in the central region. Murray & Chang (2012) explore their models by varying p from 0 (uniform), 1, and 2 (isothermal).

The density structure is important for the physical and chemical evolution of HMSFRs. It is therefore important to quantify the initial density profile on cloud to clump and cores scales and how it changes with time. While theoretical models usually do not predict, but rather assume a given density profile, observations of HMSFRs on different scales can help to narrow down the parameter space (see Fig. 10). Hydrodynamic simulations reported by Chen et al. (2020) investigate how changes of p in giant molecular clouds with an initial radius of 20 pc affect massive star cluster formation. They find that for steep density profiles, p = 2, there is a centrally-concentrated cluster, while for shallower profiles hierarchical fragmentation occurs. Hydrodynamic simulations by Girichidis et al. (2011) show that massive protostars form only in clouds with a density index of p = 1.5 or p = 2.0, while for uniform or Bonnor-Ebert-like profiles a large fraction of low-mass stars form. They found that turbulence and the initial density profile are important aspects for the evolution of the cloud and the formation of clusters.

We study correlations of all core properties shown in Table 3 using the Spearman correlation coefficient r. This statistical tool can be used to check if two data sets have a positive correlation (r = 1), negative correlation (r = -1), or no correlation (r = 0). We define that a high correlation exists if r > 0.8. For example, Feng et al. (2020) finds a negative correlation of the H₂ column density and dust temperature for cold high-mass clumps using the Spearman correlation coefficient r. A big advantage compared to the Pearson correlation coefficient is that linear, as well as nonlinear correlations are considered in the calculation of r. In addition, we add the M/L ratio of the region listed in Table 1 in Beuther et al. (2018) as a parameter for each core. However, the interpretation is difficult as multiple cores within a region have the same M/L ratio. A mean chemical age is used in the computation of r for cores for which only a time range can be estimated (Table 3).

The results for the correlation coefficient *r* are shown in Fig. 11, where all pairs with a correlation > 0.8 are highlighted. Unfortunately, a small sample of only 22 cores does not allow us to find many strong correlations. Observations of many HMSFRs at core-scales are required to study these relations in a better statistical way, which will be possible, e.g., with the ALMAGAL survey, an ongoing ALMA large program observing more than 1000 HMSFRs. A high correlation is found between the inner and outer radius, which is due to the fact that we are resolution limited and the regions are located at different distances. The correlation between p and α is due to Eq. (3). However, we find a strong negative correlation between the M/L ratio and the H₂ column density of the cores, so more evolved cores have a higher beam-convolved H₂ column density. The M/L ratio, proposed to be a good tracer of the evolutionary stage, will be investigated in the following section.

6.2. M/L ratio as a tracer of evolutionary trend

Sridharan et al. (2002) found that the distance-independent ratio M/L of UCHII regions is lower (~0.005) than the ratio of HMPOs (~0.025), as UCHII regions are more evolved and thus more luminous. This has also been confirmed by observations of large samples of HMSFRs, e.g., by Molinari et al. (2008, 2010); Maud et al. (2015); Molinari et al. (2016); Urquhart et al. (2018); Molinari et al. (2019). In Fig. 12 we plot the region-average M/Lratio taken from Table 1 in Beuther et al. (2018) against the estimated chemical ages $\tau_{\rm chem}$. The chemical timescale and a factor 2 uncertainty is shown for cores with a clear best-fit model. For cores with estimated chemical age ranges, the mean chemical age are shown with error bars spanning over the time range. The luminosities L have uncertainties on the order of $\sim 30\%$ (Mottram et al. 2011b) and the masses M calculated in Beuther et al. (2018) are expected to be uncertain within a factor of 3. We also show the average M/L ratio for HMPOs and UCHII regions derived by Sridharan et al. (2002). We corrected the M/L ratios of Sridharan et al. (2002) by a factor of 0.5, as M was taken from Beuther et al. (2002) where the reported values for *M* were lower by a factor of 2 (Beuther et al. 2005). The cores G108.75 1 and 2 are excluded here, because no consistent continuum data is available to reliably derive the mass (Beuther et al. 2018). There is a tendency such that the older cores have a lower M/L ratio. However, as the M/L ratios are determined on much larger scales and we resolve multiple cores within each region, it is difficult to compare these properties.

Estimating the chemical ages with MUSCLE shows that a linerich spectrum does not have to imply that a core is more evolved compared to a core with a line-poor spectrum. For example, the line-poor core 1 in IRAS 23033 is estimated to be older than the line-richer cores 2 and 3. When a hot core evolves to become an UCHII region, the strong protostellar radiation destroys molecules. Hence line-poor spectra can be found at early and at late evolutionary stages. Observations of more species would be helpful to better constrain the chemical model.

We are not able to derive radial temperature profiles from the temperature maps of G138.2957, G139.9091, and S106 (Fig. C.1). In G139.9091 and S106 there is no emission at the adopted $10\sigma_{\text{line,map}}$ level. There is diffuse emission of H₂CO in G138.2957, but it is too diffuse to derive a radial temperature profile. The spectra of these regions are line-poor and only simple species are detected (¹³CO, C¹⁸O, SO, H₂CO, see Fig. F.1). Based on the shape of the 1.37 mm continuum emission, G139.9091 and S106 have isolated cores with no significant envelope emission, while G138.2957 has diffuse dust emission (Fig. 3). Therefore one may conclude that G139.9091 and S106 are already more evolved and probably have $\tau_{\rm chem} > 100\,000\,{\rm yrs}$. S106 is a well studied bipolar HII region (e.g., Roberts et al. 1995; Schneider et al. 2007). G139.9091 is associated with a HII region as well (e.g., Kurtz et al. 1994; Purser 2017; Obonyo et al. 2019). G138.2957 could be in a very young strongly depleted phase with $\tau_{\rm chem}$ < 20000 yrs or could be an evolved region and the diffuse emission is due to the disruption by the protostellar radiation. With observations of G138.2957 at 5.8 cm and 20 cm, a core component with an associated synchrotron jet is seen toward the location of the 1.37 mm continuum peak and the position has an associated infrared source (Obonyo et al. 2019). This suggests that G138.2957 is an evolved embedded cluster and not a young region. Deep observations at radio wavelengths $(\sim 5-50 \text{ GHz})$ would also provide information on the presence of UCHII regions within the CORE sample. We find that for the known HII regions, we only detect simple species such as CO isotopologues, H₂CO suggesting that a large fraction of the larger molecules are already destroyed in this stage.

6.3. Correlations between chemical species

In this section, we aim to analyze which molecules show correlations by chemical links or temperature effects. We compute the Spearman correlation coefficient *r* for the molecular column density pair combinations relative to N(C¹⁸O). SO₂, HC₃N; $v_7 = 1$, and HC₃N; $v_7 = 2$ are excluded from this analysis, as there are less than 20 column density data points. Ideally, we would compare the correlations of relative abundance pairs relative to N(H₂), but due to the missing flux issue, we use N(C¹⁸O) which is also detected toward all positions and is optically thin (Table B.1). Comparing the correlation between column density pairs, one faces the issue that toward higher densities the column density is also higher as discussed in Sect. 4.2. The core and non core positions were equally considered in the calculation of *r* and the results for all column density pairs are shown in Fig. 13.

In general, all pairs show a positive correlation, r > 0. High correlation coefficients (r > 0.8) are found between pairs of the following molecules: HC₃N–SO, HC₃N–OCS, CH₃CN–



Fig. 13: Spearman correlation coefficient *r* for pairs of molecular column densities relative to N($C^{18}O$). Values higher than 0.8 are marked by a green circle.

SO, CH₃CN–OCS, CH₃CN–H₂CO, CH₃CN–HNCO, CH₃CN–HC₃N, CH₃OH–CH₃OH; $v_t = 1$, CH₃CN–CH₃OH. The lowest correlations are found for ¹³CO and DCN where they show no correlation with any species. In the case of ¹³CO this is most likely due to a high optical depth (Table B.1), so the column density cannot be reliably derived. The case of DCN is more puzzling, but a more detailed study of the deuteration would be required. Unfortunately, the CORE spectral setup covers only the DCN 3–2 line, so follow-up observations of deuterated species are necessary to study how deuterium chemistry behaves on such scales.

Methyl cyanide (CH₃CN) shows a strong positive correlation with most other species, including S-bearing and Nbearing species. A correlation of HC3N-CH3CN can be explained by gas-phase N-chemistry in the envelope gas (Bergner et al. 2017). We find that OCS is correlated with other dense gas tracers (HC₃N and CH₃CN). A correlation exists between CH₃OH and CH₃CN even though, there is no chemical link between methanol and methyl cyanide. Such a correlation has also been found toward low-mass star-forming regions (Bergner et al. 2017; Belloche et al. 2020) and toward the massive star-forming region G10.6-0.4 (Law et al. 2021). Belloche et al. (2020) argued that this is a temperature effect caused by chemically unrelated species being evaporated from icy grain mantles by energetic processes. Additional high angular resolution observations of SiO would be helpful to study the impact of shocks in more detail. In the 1D physical-chemical modeling of HMSFRs with MUSCLE by Gerner et al. (2014), CH₃CN and CH₃OH are cospatial in radial abundance profiles toward the inner hot core region. Urquhart et al. (2019) find that the line integrated ratios of S- and N-bearing species are positively correlated with the dust temperature in a sample of high-mass star-forming clumps. Gratier et al. (2013) found that CH₃CN is much more abundant in the photo-dissociation region (PDR) of the Horsehead nebula than the associated cold and dense core. This is consistent with the fact that Purcell et al. (2006) detect 3 mm lines of CH₃CN in 58 candidate hot molecular cores on a sample of 83 methanol maser-selected star-forming regions. They detect CH₃CN in isolated methanol maser sites and find that CH₃CN is more prevalent and brighter when an UCHII region is present, independent of the distance to the source. Guzmán et al. (2014) propose that



Fig. 14: Comparison of the free-fall timescale $\tau_{\rm ff}$ and crossing timescale $\tau_{\rm cross}$ (colored lines), and chemical timescales $\tau_{\rm chem}$ (black data points).

correlated abundances of CH₃OH and CH₃CN could be related with photochemistry, e.g., by photodesorption.

The column densities relative to $N(C^{18}O)$ show in general positive correlations. N-bearing and S-bearing species seem to be chemically related by high temperature gas-phase chemistry. The strong non-correlation of DCN with any observed species needs high angular resolution follow-up observations of deuterated molecules toward these HMSFRs (e.g., N2D⁺, DCO⁺). The correlation between CH₃CN and CH₃OH can be due to a mutual evaporation temperature and/or due to photo-desorption.

Computing the Spearman correlation coefficient r of the chemical timescales τ_{chem} (Table 3) with the observed column densities N (Fig. 8) and with the relative abundances $N/N(C^{18}O)$ (Fig. D.1) for all 22 cores, we do not find strong correlations for most of the molecular column densities and relative abundances with the chemical age. We use the mean chemical age for cores for which only a time range can be estimated. Positive correlations of the column density and relative abundance with the chemical timescale (r = 0.5 - 0.9) are found for SO₂, HNCO, $HC_3N;v_7 = 1$, and $CH_3OH;v_t = 1$. We only detect $HC_3N;v_7 = 2$ in cores which have a chemical age estimated to be $> 80\,000$ yrs. This suggests that vibrationally and torsionally excited states of molecules are good indicators for a more evolved region, as these transitions have higher upper energy levels (Table B.1) and are only excited at a high kinetic temperature. We propose that SO₂ and HNCO are also good tracers of the evolutionary stage, however, this has to be investigated in a larger statistical sample.

6.4. Comparison of physical and chemical timescales

In the following, we compare the estimated chemical timescales with commonly applied physical timescales. The free-fall timescale $\tau_{\rm ff}$ is the time it takes for a spherical object to fully collapse under the influence of gravity with no additional forces.

It only depends on the initial density ρ :

$$\tau_{\rm ff} = \sqrt{\frac{3\pi}{32G\rho(\rm H_2)}},\tag{7}$$

where G is the gravitational constant and $\rho(H_2)$ the initial H₂ mass density.

The crossing timescale is the time it takes to cross the system once:

$$\tau_{\rm cross} = \frac{R}{v},\tag{8}$$

with clump radius R and velocity dispersion v.

A comparison between the free-fall timescales for corresponding H_2 number densities of 10^5 cm⁻³, 10^6 cm⁻³, 10^7 cm⁻³, and 10^8 cm^{-3} , crossing timescale (assuming R = 1 pc and v =4 km s⁻¹, Elmegreen 2000) and the chemical ages of the regions estimated with MUSCLE is shown in Fig. 14. A mean chemical age is used for cores for which only a time range can be determined. We then calculate the mean chemical age for regions associated with multiple cores. In these cases, the errorbars indicate the range of chemical ages. For regions associated with only one core, we assume a factor of 2 uncertainty for cores with a clear best-fit chemical age or the errorbars indicate the estimated time range for cores for which we can only determine a time range. The derived chemical ages are in agreement with an initial clump density of $10^5 - 10^6$ cm⁻³. Indeed, observations suggest that IRDCs have densities $n(H_2) > 10^5 \text{ cm}^{-3}$ (e.g., Carey et al. 1998). The estimated crossing time is an upper limit, in agreement with the scenario that suggests that star formation occurs within a few crossing times (Elmegreen 2000). The agreement of the physical timescales with the chemical ages shows that even though the physical-chemical model is pseudo time-dependent, the estimates are feasible, especially when a homogeneous data set and analysis is used.

Other forces such as magnetic fields and turbulence can slow down or prevent collapse and are important in the inital diffuse ISM (Ballesteros-Paredes et al. 2007; Burge et al. 2016). However, in the diffuse ISM, molecules, with the exception of H₂, have not formed yet. Our zero point for the chemical timescales is when the density reaches 10^4 cm^{-3} and for the freefall timescale we also assume densities $\ge 10^4 \text{ cm}^{-3}$. In Beuther et al. (2018), we showed that the fragmentation scales we observe is in agreement with thermal Jeans fragmentation and turbulent Jeans fragmentation is not important. This is fully consistent with the results found in a different sample by Palau et al. (2015). The role of magnetic fields in the CORE regions are being investigated by polarization observations with the Submillimeter Array (SMA).

7. Conclusions

We study the physical and chemical structure of 18 high-mass star-forming regions using the CORE 1 mm continuum and spectral line data. We quantify the chemical content for in total 120 positions including in total 22 cores. Combining the CORE observations with the physical-chemical model MUSCLE, we estimate the chemical timescales of the cores. Our main conclusions are summarized as follows:

1. Using H₂CO and CH₃CN line emission, we derive temperature maps of the regions. We identify "cores" as objects having a clear radially decreasing temperature profile. We fit these radial temperature profiles and obtain an average power-law index $q = 0.4 \pm 0.1$ excluding three outliers with $q \ge 1$, which is in agreement with theoretical predictions and calculations.

- 2. The 1.37 mm continuum visibility profiles of the cores reveal a mean density power-law index of $p = 2.0 \pm 0.2$. Comparing these high-resolution density profiles to previous singledish and interferometric studies, the density profiles appear to stay roughly constant between p = 1.7 - 2.0 from scales of 1 pc down to 1 000 au. The molecular hydrogen column density $N(H_2)$ has a mean of 1.5×10^{24} cm⁻² toward the 120 positions and the core mass $M_{\rm core}$ of the 22 cores has a mean of $4.1 M_{\odot}$.
- 3. We derive molecular column densities of 11 species by fitting their spectral lines with XCLASS. Spearman correlation coefficients are evaluated for all molecule pairs. We find high correlations between N-bearing and S-bearing species that are chemically related through high temperature gas-phase chemistry. High correlations are also found for molecules that are not chemically related (CH₃CN and CH₃OH), but for which the correlation results from a common evaporation temperature and/or photo-desorption.
- 4. We apply the physical-chemical model MUSCLE to the observed column densities of each core to estimate the chemical age τ_{chem} . We find a spread in age from $\tau_{chem} \sim 20\,000 100\,000$ yrs and a mean chemical age of $\sim 60\,000$ yrs. Multiple cores within a region show that there can be an age gradient for largely separated cores (e.g. toward IRAS 23033 and G108.75), while close cores have a similar chemical age (e.g. toward IRAS 21078 and W3 H2O) suggesting a sequential star formation.
- 5. A strong correlation is found between the peak column density $N(H_2)$ of the 22 cores and the M/L ratio of the region. In addition, we find a trend that older cores have a lower M/Lratio. However, a larger sample is required to study the physical and chemical properties on core-scales in more detail.
- 6. We compare the chemical age with physical timescales. We find that the chemical age is consistent with a free-fall timescale of the initial clump at a density of 10⁵ cm⁻³ and 10⁶ cm⁻³ consistent with density values typically found toward IRDCs. The chemical ages are smaller than a crossing time of the parental clump.
- 7. We improve the quality (both intensity and noise) of the NOEMA continuum and spectral line data by applying self-calibration. All of the standard and self-calibrated CORE data are publically available at https://www.mpia.de/core.

The CORE data reveal a large physical and chemical diversity on scales down to \sim 300 au. Here, we use the molecular column densities of 120 positions toward the 18 high-mass star-forming regions to quantify the chemical content, with an emphasis on 22 cores. The case-study of AFGL 2591 already revealed complex structures around a single hot core (Gieser et al. 2019) using the high resolution interferometric observations. In a next step, we will study the spatial morphology of the molecular emission in order to study spatial correlations of the species.

Acknowledgements. The authors would like to thank the anonymous referee whose comments helped improve the clarity of this paper. This work is based on observations carried out under project number L14AB with the IRAM NOEMA Interferometer and the IRAM 30m telescope. IRAM is supported by INSU/CNRS (France), MPG (Germany), and IGN (Spain). We thank the IRAM staff, especially R. Neri and S. Bardeau, for their help in obtaining, calibrating, and imaging the NOEMA and IRAM 30m data. C. G., H. B., and S. S. acknowledge support from the European Research Council under the Horizon

2020 Framework Programme via the ERC Consolidator Grant CSF-648505. D. S. acknowledges support by the Deutsche Forschungsgemeinschaft through SPP 1833: "Building a Habitable Earth (SE 1962/6-1)". R. G.-M. acknowledges support from UNAM-PAPIIT project IN104319. R. K. acknowledges financial support via the Emmy Noether Research Group on Accretion Flows and Feedback in Realistic Models of Massive Star Formation funded by the German Research Foundation (DFG) under grant no. KU 2849/3-1 and KU 2849/3-2. A. S.-M. research is partially carried out within the Collaborative Research Centre 956 (subproject A6), funded by the Deutsche Forschungsgemeinschaft (DFG) - project ID 184018867. A. P. acknowledges financial support from CONACyT and UNAM-PAPIIT IN113119 grant, México. J. P. acknowledges support by the Programme National "Physique et Chimie du Milieu Interstellaire" (PCMI) of CNRS/INSU with INC/INP co-funded by CEA and CNES. This research made use of Astropy³, a community-developed core Python package for Astronomy (Astropy Collaboration et al. 2013, 2018).

References

- Adams, F. C. 1991, ApJ, 382, 544
- Ahmadi, A., Beuther, H., Mottram, J. C., et al. 2018, A&A, 618, A46
- Ahmadi, A., Kuiper, R., & Beuther, H. 2019, A&A, 632, A50
- Allen, V., van der Tak, F. F. S., Sánchez-Monge, Á., Cesaroni, R., & Beltrán, M. T. 2017, A&A, 603, A133
- Astropy Collaboration, Price-Whelan, A. M., Sipőcz, B. M., et al. 2018, AJ, 156, 123
- Astropy Collaboration, Robitaille, T. P., Tollerud, E. J., et al. 2013, A&A, 558, A33
- Ballesteros-Paredes, J., Klessen, R. S., Mac Low, M. M., & Vazquez-Semadeni, E. 2007, in Protostars and Planets V, ed. B. Reipurth, D. Jewitt, & K. Keil, 63
- Belloche, A., Maury, A. J., Maret, S., et al. 2020, A&A, 635, A198
- Belloche, A., Müller, H. S. P., Menten, K. M., Schilke, P., & Comito, C. 2013, A&A, 559, A47
- Beltrán, M. T. & de Wit, W. J. 2016, A&A Rev., 24, 6
- Bergner, J. B., Öberg, K. I., Garrod, R. T., & Graninger, D. M. 2017, ApJ, 841, 120
- Bertin, M., Romanzin, C., Doronin, M., et al. 2016, ApJ, 817, L12
- Beuther, H., Churchwell, E. B., McKee, C. F., & Tan, J. C. 2007a, Protostars and Planets V, 165
- Beuther, H., Henning, T., Linz, H., et al. 2010, A&A, 518, L78
- Beuther, H., Leurini, S., Schilke, P., et al. 2007b, A&A, 466, 1065
- Beuther, H., Mottram, J. C., Ahmadi, A., et al. 2018, A&A, 617, A100
- Beuther, H., Schilke, P., Menten, K. M., et al. 2002, ApJ, 566, 945
- Beuther, H., Schilke, P., Menten, K. M., et al. 2005, ApJ, 633, 535
- Bonfand, M., Belloche, A., Menten, K. M., Garrod, R. T., & Müller, H. S. P. 2017, A&A, 604, A60
- Bonnell, I. A. 2007, in Astronomical Society of the Pacific Conference Series, Vol. 367, Massive Stars in Interactive Binaries, ed. N. St.-Louis & A. F. J. Moffat, 303
- Bonnell, I. A. & Bate, M. R. 2002, MNRAS, 336, 659
- Bonnell, I. A., Bate, M. R., Clarke, C. J., & Pringle, J. E. 2001, MNRAS, 323, 785
- Bonnell, I. A., Bate, M. R., & Zinnecker, H. 1998, MNRAS, 298, 93
- Bosco, F., Beuther, H., Ahmadi, A., et al. 2019, A&A, 629, A10
- Bottinelli, S., Ceccarelli, C., Neri, R., et al. 2004, ApJ, 617, L69
- Burge, C. A., Van Loo, S., Falle, S. A. E. G., & Hartquist, T. W. 2016, A&A, 596, A28
- Butler, M. J. & Tan, J. C. 2012, ApJ, 754, 5
- Carey, S. J., Clark, F. O., Egan, M. P., et al. 1998, ApJ, 508, 721
- Carter, M., Lazareff, B., Maier, D., et al. 2012, A&A, 538, A89
- Caselli, P., Hasegawa, T. I., & Herbst, E. 1993, ApJ, 408, 548
- Cesaroni, R., Beuther, H., Ahmadi, A., et al. 2019, A&A, 627, A68
- Cesaroni, R., Felli, M., Testi, L., Walmsley, C. M., & Olmi, L. 1997, A&A, 325, 725
- Chen, Y., Li, H., & Vogelsberger, M. 2020, arXiv e-prints, arXiv:2006.00004
- Churchwell, E. 2002, ARA&A, 40, 27
- Clark, B. G. 1980, A&A, 89, 377
- Cooper, H. D. B. 2013, PhD thesis, University of Leeds
- Cruz-Diaz, G. A., Martín-Doménech, R., Muñoz Caro, G. M., & Chen, Y.-J. 2016, A&A, 592, A68
- Cuppen, H. M., Walsh, C., Lamberts, T., et al. 2017, Space Sci. Rev., 212, 1
- Draine, B. T. 2011, Physics of the Interstellar and Intergalactic Medium (Princeton: Princeton University Press)
- Duarte-Cabral, A., Bontemps, S., Motte, F., et al. 2013, A&A, 558, A125 Elmegreen, B. G. 2000, ApJ, 530, 277
- ³ http://www.astropy.org

Article number, page 28 of 81

- Emerson, J. P. 1988, in NATO Advanced Science Institutes (ASI) Series C, Vol. 241, NATO Advanced Science Institutes (ASI) Series C, ed. A. K. Dupree & M. T. V. T. Lago, 21
- Endres, C. P., Schlemmer, S., Schilke, P., Stutzki, J., & Müller, H. S. P. 2016, Journal of Molecular Spectroscopy, 327, 95
- Fayolle, E. C., Öberg, K. I., Garrod, R. T., van Dishoeck, E. F., & Bisschop, S. E. 2015, A&A, 576, A45
- Feng, S., Beuther, H., Henning, T., et al. 2015, A&A, 581, A71
- Feng, S., Beuther, H., Semenov, D., et al. 2016, A&A, 593, A46
- Feng, S., Li, D., Caselli, P., et al. 2020, ApJ, 901, 145
- Frau, P., Girart, J. M., Beltrán, M. T., et al. 2010, ApJ, 723, 1665
- Gerin, M. 2013, The Molecular Universe, ed. I. W. M. Smith, C. S. Cockell, & S. Leach (Berlin, Heidelberg: Springer Berlin Heidelberg), 35-72
- Gerner, T., Beuther, H., Semenov, D., et al. 2014, A&A, 563, A97
- Gerner, T., Shirley, Y. L., Beuther, H., et al. 2015, A&A, 579, A80
- Giannetti, A., Brand, J., Sánchez-Monge, Á., et al. 2013, A&A, 556, A16
- Gieser, C., Semenov, D., Beuther, H., et al. 2019, A&A, 631, A142
- Girichidis, P., Federrath, C., Banerjee, R., & Klessen, R. S. 2011, MNRAS, 413, 2741
- Gratier, P., Pety, J., Guzmán, V., et al. 2013, A&A, 557, A101
- Guzmán, V. V., Pety, J., Gratier, P., et al. 2014, Faraday Discussions, 168, 103
- Hartmann, L., Ballesteros-Paredes, J., & Heitsch, F. 2012, MNRAS, 420, 1457
- Hatchell, J., Thompson, M. A., Millar, T. J., & MacDonald, G. H. 1998, A&AS, 133, 29
- Hatchell, J. & van der Tak, F. F. S. 2003, A&A, 409, 589
- Herbst, E. & van Dishoeck, E. F. 2009, ARA&A, 47, 427
- Hildebrand, R. H. 1983, QJRAS, 24, 267
- Indriolo, N., Neufeld, D. A., Gerin, M., et al. 2015, ApJ, 800, 40
- Jiménez-Serra, I., Zhang, Q., Viti, S., Martín-Pintado, J., & de Wit, W. J. 2012, ApJ, 753, 34
- Johnston, K. G., Hoare, M. G., Beuther, H., et al. 2020, ApJ, 896, 35
- Jørgensen, J. K., Belloche, A., & Garrod, R. T. 2020, ARA&A, 58, 727
- Kauffmann, J., Bertoldi, F., Bourke, T. L., Evans, N. J., I., & Lee, C. W. 2008, A&A, 487, 993
- Kennicutt, Robert C., J. 1998, ARA&A, 36, 189
- Krumholz, M. R. 2015, Astrophysics and Space Science Library, Vol. 412, The Formation of Very Massive Stars, ed. J. S. Vink, 43
- Kuiper, R. & Hosokawa, T. 2018, A&A, 616, A101 Kurtz, S., Churchwell, E., & Wood, D. O. S. 1994, ApJS, 91, 659
- Larson, R. B. 1981, MNRAS, 194, 809
- Law, C. J., Zhang, Q., Öberg, K. I., et al. 2021, arXiv e-prints, arXiv:2101.07801
- Lee, H. H., Roueff, E., Pineau des Forets, G., et al. 1998, A&A, 334, 1047
- Looney, L. W., Mundy, L. G., & Welch, W. J. 2003, ApJ, 592, 255
- Mangum, J. G. & Shirley, Y. L. 2015, PASP, 127, 266
- Mangum, J. G. & Wootten, A. 1993, ApJS, 89, 123
- Maud, L. T., Lumsden, S. L., Moore, T. J. T., et al. 2015, MNRAS, 452, 637 McGuire, B. A. 2018, ApJS, 239, 17 McKee, C. F. & Ostriker, E. C. 2007, ARA&A, 45, 565

- McKee, C. F. & Tan, J. C. 2002, Nature, 416, 59 McKee, C. F. & Tan, J. C. 2003, ApJ, 585, 850
- McLaughlin, D. E. & Pudritz, R. E. 1996, ApJ, 469, 194
- McLaughlin, D. E. & Pudritz, R. E. 1997, ApJ, 476, 750
- Mills, E. A. C., Corby, J., Clements, A. R., et al. 2018, ApJ, 869, 121
- Molinari, S., Baldeschi, A., Robitaille, T. P., et al. 2019, MNRAS, 486, 4508
- Molinari, S., Merello, M., Elia, D., et al. 2016, ApJ, 826, L8
- Molinari, S., Pezzuto, S., Cesaroni, R., et al. 2008, A&A, 481, 345 Molinari, S., Swinyard, B., Bally, J., et al. 2010, PASP, 122, 314
- Möller, T., Endres, C., & Schilke, P. 2017, A&A, 598, A7
- Moscadelli, L., Beuther, H., Ahmadi, A., et al. 2021, arXiv e-prints, arXiv:2102.04872
- Motte, F. & André, P. 2001, A&A, 365, 440

Molecular Structure, 742, 215

453, 3785

- Motte, F., Bontemps, S., & Louvet, F. 2018, Annual Review of Astronomy and Astrophysics, 56, 41
- Mottram, J. C., Beuther, H., Ahmadi, A., et al. 2020, A&A, 636, A118
- Mottram, J. C., Hoare, M. G., Davies, B., et al. 2011a, ApJ, 730, L33 Mottram, J. C., Hoare, M. G., Urquhart, J. S., et al. 2011b, A&A, 525, A149
- Mueller, K. E., Shirley, Y. L., Evans, Neal J., I., & Jacobson, H. R. 2002, ApJS, 143, 469 Müller, H. S. P., Schlöder, F., Stutzki, J., & Winnewisser, G. 2005, Journal of

Murray, N. & Chang, P. 2012, ApJ, 746, 75 Obonyo, W. O., Lumsden, S. L., Hoare, M. G., et al. 2019, MNRAS, 486, 3664

Palau, A., Ballesteros-Paredes, J., Vázquez-Semadeni, E., et al. 2015, MNRAS,

Palau, A., Zhang, Q., Girart, J. M., et al. 2020, arXiv e-prints, arXiv:2010.12099

Olguin, F. A., Hoare, M. G., Johnston, K. G., et al. 2020, MNRAS, 498, 4721

Osorio, M., Lizano, S., & D'Alessio, P. 1999, ApJ, 525, 808

Palau, A., Estalella, R., Girart, J. M., et al. 2014, ApJ, 785, 42

Pearson, T. J. & Readhead, A. C. S. 1984, ARA&A, 22, 97

Ossenkopf, V. & Henning, T. 1994, A&A, 291, 943

- Peters, T., Mac Low, M.-M., Banerjee, R., Klessen, R. S., & Dullemond, C. P. 2010, ApJ, 719, 831
- Pickett, H. M., Poynter, R. L., Cohen, E. A., et al. 1998, J. Quant. Spectr. Rad. Transf., 60, 883
- Pillai, T., Wyrowski, F., Carey, S. J., & Menten, K. M. 2006, A&A, 450, 569
- Purcell, C. R., Balasubramanyam, R., Burton, M. G., et al. 2006, MNRAS, 367, 553
- Purser, S. J. D. 2017, PhD thesis, University of Leeds
- Qin, S.-L., Schilke, P., Wu, J., et al. 2015, ApJ, 803, 39
- Radcliffe, J. F., Garrett, M. A., Beswick, R. J., et al. 2016, A&A, 587, A85
- Rathborne, J. M., Jackson, J. M., & Simon, R. 2006, ApJ, 641, 389
- Remijan, A. J., Markwick-Kemper, A., & ALMA Working Group on Spectral Line Frequencies. 2007, in American Astronomical Society Meeting Abstracts, Vol. 211, American Astronomical Society Meeting Abstracts, 132.11
- Roberts, D. A., Crutcher, R. M., & Troland, T. H. 1995, ApJ, 442, 208
- Rodgers, S. D. & Charnley, S. B. 2001, ApJ, 546, 324
- Rodón, J. A., Beuther, H., & Schilke, P. 2012, A&A, 545, A51
- Rosen, A. L., Offner, S. S. R., Sadavoy, S. I., et al. 2020, Space Sci. Rev., 216, 62
- Sánchez-Monge, Á., Cesaroni, R., Beltrán, M. T., et al. 2013a, A&A, 552, L10
- Sánchez-Monge, Á., Kurtz, S., Palau, A., et al. 2013b, ApJ, 766, 114
- Sault, R. J., Teuben, P. J., & Wright, M. C. H. 1995, in Astronomical Society of the Pacific Conference Series, Vol. 77, Astronomical Data Analysis Software and Systems IV, ed. R. A. Shaw, H. E. Payne, & J. J. E. Hayes, 433
- Schilke, P. 2015, in EAS Publications Series, Vol. 75, EAS Publications Series, 227–235
- Schilke, P., Walmsley, C. M., Pineau des Forets, G., & Flower, D. R. 1997, A&A, 321, 293
- Schneider, N., Simon, R., Bontemps, S., Comerón, F., & Motte, F. 2007, A&A, 474, 873
- Schöier, F. L., van der Tak, F. F. S., van Dishoeck, E. F., & Black, J. H. 2005, A&A, 432, 369
- Semenov, D., Hersant, F., Wakelam, V., et al. 2010, A&A, 522, A42
- Shu, F. H. 1977, ApJ, 214, 488
- Shu, F. H., Adams, F. C., & Lizano, S. 1987, ARA&A, 25, 23
- Smith, R. J., Longmore, S., & Bonnell, I. 2009, MNRAS, 400, 1775
- Sridharan, T. K., Beuther, H., Schilke, P., Menten, K. M., & Wyrowski, F. 2002, ApJ, 566, 931
- Steer, D. G., Dewdney, P. E., & Ito, M. R. 1984, A&A, 137, 159
- Tan, J. C., Beltrán, M. T., Caselli, P., et al. 2014, Protostars and Planets VI, 149
- Urquhart, J. S., Figura, C., Wyrowski, F., et al. 2019, MNRAS, 484, 4444
- Urquhart, J. S., König, C., Giannetti, A., et al. 2018, MNRAS, 473, 1059
- van der Tak, F. F. S., van Dishoeck, E. F., Evans, Neal J., I., & Blake, G. A. 2000, ApJ, 537, 283
- Williams, J. P., de Geus, E. J., & Blitz, L. 1994, ApJ, 428, 693
- Williams, S. J., Fuller, G. A., & Sridharan, T. K. 2004, A&A, 417, 115
- Williams, S. J., Fuller, G. A., & Sridharan, T. K. 2005, A&A, 434, 257
- Wilson, T. L. & Rood, R. 1994, ARA&A, 32, 191
- Wood, D. O. S. & Churchwell, E. 1989, ApJS, 69, 831
- Wyrowski, F., Güsten, R., Menten, K. M., et al. 2016, A&A, 585, A149
- Wyrowski, F., Schilke, P., Walmsley, C. M., & Menten, K. M. 1999, ApJ, 514, L43
- Zhang, Q., Wang, Y., Pillai, T., & Rathborne, J. 2009, ApJ, 696, 268
- Zinnecker, H. & Yorke, H. W. 2007, ARA&A, 45, 481

- ¹ Max Planck Institute for Astronomy, Königstuhl 17, 69117 Heidelberg, Germany, e-mail: gieser@mpia.de
- ² Department of Chemistry, Ludwig Maximilian University, Butenandtstr. 5-13, 81377 Munich, Germany
- ³ Leiden University, Niels Bohrweg 2, 2333 CA Leiden, Netherlands
- ⁴ I. Physikalisches Institut, Universität zu Köln, Zülpicher Str. 77, D-50937, Köln, Germany
- ⁵ INAF, Osservatorio Astrofisico di Arcetri, Largo E. Fermi 5, I-50125 Firenze, Italy
- ⁶ UK Astronomy Technology Centre, Royal Observatory Edinburgh, Blackford Hill, Edinburgh EH9 3HJ, UK
- ⁷ Center for Astrophysics | Harvard & Smithsonian, 60 Garden Street, Cambridge, MA 02138, USA
- ⁸ Centre for Astrophysics and Planetary Science, University of Kent, Canterbury, CT2 7NH, UK
- ⁹ Academia Sinica Institute of Astronomy and Astrophysics, No.1, Sec. 4, Roosevelt Rd, Taipei 10617, Taiwan, Republic of China
- ¹⁰ CAS Key Laboratory of FAST, National Astronomical Observatories, Chinese Academy of Sciences, Beijing 100101, People's Republic of China
- ¹¹ National Astronomical Observatory of Japan, National Institutes of Natural Sciences, 2-21-1 Osawa, Mitaka, Tokyo 181-8588, Japan
- ¹² Instituto de Radioastronomía y Astrofísica (IRyA), UNAM, Apdo. Postal 72-3 (Xangari), Morelia, Michoacán 58089, Mexico
- ¹³ Institut de Radioastronomie Millimétrique (IRAM), 300 Rue de la Piscine, F-38406 Saint Martin d'Hères, France
- ¹⁴ Astrophysics Research Institute, Liverpool John Moores University, Liverpool, L3 5RF, UK
- ¹⁵ INAF, Osservatorio Astronomico di Cagliari, Via della Scienza 5, I-09047, Selargius (CA), Italy
- ¹⁶ Institute of Astronomy and Astrophysics, University of Tübingen, Auf der Morgenstelle 10, 72076, Tübingen, Germany
- ¹⁷ Max-Planck-Institut für Astrophysik, Karl-Schwarzschild-Str. 1, 85748 Garching, Germany
- ¹⁸ Max Planck Institut for Radioastronomie, Auf dem Hügel 69, 53121 Bonn, Germany
- ¹⁹ Laboratoire d'astrophysique de Bordeaux, Univ. Bordeaux, CNRS, B18N, allèe Geoffroy Saint-Hilaire, 33615 Pessac, France
- ²⁰ Physics Department, UMIST, P.O. Box 88, Manchester M60 1QD, UK
- ²¹ School of Physics and Astronomy, University of Leeds, Leeds LS2 9JT, United Kingdom
- ²² European Southern Observatory, Karl-Schwarzschild-Str. 2, D-85748 Garching, Germany
- ²³ LERMA, Observatoire de Paris, PSL Research University, CNRS, Sorbonne Universités, 75014 Paris, France
- ²⁴ Department of Physics and Astronomy, McMaster University, 1280 Main St. W, Hamilton, ON L8S 4M1, Canada

Appendix A: Position properties

Table A.1 summarizes the properties of all 120 positions extracted within the 18 CORE regions which were analyzed in this study. The positions are also marked in Fig. 3 showing the 1.37 mm continuum emission. Positions which show a clear temperature profile are labeled as "C" (further explained in Sect. 3.2).

Table A.1: Overview of selected positions in each region. The systemic velocity is derived from the C¹⁸O 2 – 1 line (Sect. 4.2). The noise of the spectrum σ_{line} is computed in a line-free range from a rest-frequency of 219.00 GHz to 219.13 GHz. The continuum optical depth τ_{ν}^{cont} is calculated according to Eq. (6) and the H₂ column density $N(\text{H}_2)$ is calculated according to Eq. (4) using the continuum intensity $I_{1.37\text{mm}}$ and T_{kin} (Sect. 4.1). As the H₂ column density is derived from the 1.37 mm dust continuum emission and we do not have short-spacing information this should be considered as an lower limit due to potential missing flux.

Position	α	δ	VLSR	$\sigma_{\rm line}$	T_{kin}	<i>I</i> _{1 37mm}	τ_{u}^{cont}	$N(\mathrm{H}_2)$
	J(2000)	J(2000)	$(\mathrm{kms^{-1}})$	(K)	(K)	$(mJv beam^{-1})$	V	(cm^{-2})
IRAS 23033 1 C	23:05:24.63	+60:08:09.2	-53.6	0.30	114.9± 8.2	38.13	5.5(-2)	$1.9(24)\pm4.0(23)$
IRAS 23033 2 C	23:05:25.04	+60:08:15.8	-53.4	0.20	167.2 ± 6.6	32.55	3.1(-2)	$1.1(24)\pm 2.2(23)$
IRAS 23033 3 C	23:05:24.92	+60:08:13.9	-54.6	0.20	160.8 ± 8.0	27.04	2.7(-2)	$9.5(23)\pm 2.0(23)$
IRAS 23033 4	23:05:24.97	+60:08:15.0	-53.7	0.19	84.9 ± 6.7	7.14	1.4(-2)	$4.9(23)\pm1.1(23)$
IRAS 23033 5	23:05:25.03	+60:08:17.3	-52.9	0.21	15.3 ± 2.5	6.30	9.5(-2)	$3.2(24) \pm 9.7(23)$
IRAS 23151 1 C	23:17:20.89	+59:28:47.6	-54.5	0.22	129.2 ± 4.7	33.78	4.3(-2)	$1.5(24)\pm3.1(23)$
IRAS 23151 2	23:17:20.84	+59:28:47.0	-54.2	0.22	168.8 ± 78.9	3.06	2.9(-3)	$1.0(23)\pm5.4(22)$
IRAS 23151 3	23:17:20.82	+59:28:48.4	-56.2	0.23	171.7 ± 42.2	2.96	2.8(-3)	$9.8(22) \pm 3.2(22)$
IRAS 23151 4	23:17:20.88	+59:28:49.0	-56.4	0.33	204.1±35.5	2.35	1.8(-3)	$6.5(22) \pm 1.7(22)$
IRAS 23151 5	23:17:20.96	+59:28:46.2	-53.7	0.29	69.0 ± 37.7	1.87	4.5(-3)	$1.6(23) \pm 1.0(23)$
IRAS 23385 1 C	23:40:54.51	+61:10:28.0	-50.9	0.20	239.5±12.1	17.84	9.3(-3)	$3.3(23)\pm6.8(22)$
IRAS 23385 2 C	23:40:54.72	+61:10:28.5	-51.3	0.25	226.0 ± 2.3	14.36	7.9(-3)	$2.8(23)\pm 5.6(22)$
IRAS 23385 3	23:40:54.44	+61:10:27.7	-50.8	0.17	72.4±37.5	13.79	2.5(-2)	$8.9(23)\pm5.2(23)$
IRAS 23385 4	23:40:54.42	+61:10:27.0	-49.9	0.21	21.5 ± 4.7	4.12	3.0(-2)	$1.1(24) \pm 3.7(23)$
IRAS 23385 5	23:40:54.32	+61:10:28.2	-50.1	0.22	262.2 ± 11.6	3.99	1.9(-3)	$6.7(22) \pm 1.4(22)$
IRAS 23385 6	23:40:54.59	+61:10:27.3	-51.0	0.18	51.1±32.1	3.28	8.7(-3)	$3.1(23)\pm2.2(23)$
IRAS 23385 7	23:40:54.49	+61:10:29.4	-50.7	0.20	164.2 ± 42.2	2.49	1.9(-3)	$6.8(22)\pm2.2(22)$
IRAS 23385 8	23:40:54.34	+61:10:27.2	-50.3	0.21	116.4±36.6	0.92	1.0(-3)	$3.6(22)\pm1.4(22)$
AFGL 2591 1 C	20:29:24.88	+40:11:19.4	- 5.0	0.47	159.9 ± 6.8	85.10	8.6(-2)	$2.9(24)\pm6.0(23)$
AFGL 2591 2	20:29:24.47	+40:11:24.0	- 7.2	0.36	39.3 ± 6.4	5.76	2.5(-2)	$9.0(23)\pm2.4(23)$
AFGL 2591 3	20:29:25.03	+40:11:21.3	- 5.7	0.40	61.6 ± 14.8	5.68	1.5(-2)	$5.4(23)\pm1.8(23)$
AFGL 2591 4	20:29:24.80	+40:11:20.5	- 5.8	0.35	94.3 ± 20.4	5.01	8.4(-3)	$3.0(23)\pm9.1(22)$
CepA HW2 1 C	22:56:17.98	+62:01:49.6	- 7.7	0.97	238.2±11.4	448.50	3.4(-1)	$1.0(25)\pm2.1(24)$
CepA HW2 2 C	22:56:17.96	+62:01:46.2	- 9.8	0.39	170.5±13.3	84.21	8.0(-2)	$2.7(24)\pm 5.9(23)$
CepA HW2 3	22:56:17.87	+62:01:50.3	- 7.4	0.48	124.9 ± 8.7	28.63	3.7(-2)	$1.3(24) \pm 2.7(23)$
CepA HW2 4	22:56:17.35	+62:01:54.8	-11.1	0.44	20.0 ± 10.0	20.45	2.3(-1)	$7.2(24) \pm 4.9(24)$
CepA HW2 5	22:56:17.89	+62:01:48.1	- 9.4	0.42	273.8 ± 8.5	14.19	8.0(-3)	$2.8(23)\pm 5.7(22)$
G084.9505 1 C	20:55:32.50	+44:06:10.2	-34.3	0.15	169.0± 0.8	5.19	5.0(-3)	$1.8(23)\pm3.5(22)$
G084.9505 2	20:55:32.22	+44:06:07.9	-34.5	0.17	81.8 ± 3.6	3.38	6.9(-3)	$2.4(23)\pm5.0(22)$
G084.9505 3	20:55:32.13	+44:06:08.8	-34.2	0.14	77.0 ± 22.5	2.18	4.7(-3)	$1.7(23)\pm6.2(22)$
G084.9505 4	20:55:32.30	+44:06:08.0	-34.0	0.15	111.2 ± 36.4	2.05	3.0(-3)	$1.1(23) \pm 4.3(22)$
G084.9505 5	20:55:32.41	+44:06:08.5	-34.2	0.14	53.7 ± 42.5	1.68	5.4(-3)	$1.9(23) \pm 1.7(23)$
G084.9505 6	20:55:32.16	+44:06:09.2	-34.2	0.17	111.2 ± 37.9	1.27	1.9(-3)	$6.7(22) \pm 2.7(22)$
G084.9505 7	20:55:32.31	+44:06:09.0	-34.7	0.17	33.3 ± 4.2	1.15	6.4(-3)	$2.3(23)\pm 5.6(22)$
G084.9505 8	20:55:32.38	+44:06:06.6	-33.9	0.17	57.2 ± 27.2	1.04	3.1(-3)	$1.1(23)\pm6.2(22)$
G094.6028 1 C	21:39:58.27	+50:14:21.0	+29.0	0.26	183.8 ± 45.4	12.84	1.2(-2)	$4.2(23)\pm1.3(23)$
G094.6028 2	21:39:58.16	+50:14:20.5	+30.0	0.18	39.1±17.7	2.18	1.0(-2)	$3.7(23) \pm 2.1(23)$
G094.6028 3	21:39:58.11	+50:14:26.1	+28.4	0.25	50.9 ± 0.0	2.11	7.5(-3)	$2.7(23)\pm 5.3(22)$
G094.6028 4	21:39:58.31	+50:14:19.6	+28.3	0.19	37.8 ± 8.5	2.05	1.0(-2)	$3.6(23)\pm1.2(23)$
G094.6028 5	21:39:58.13	+50:14:21.6	+29.0	0.17	21.7 ± 5.0	2.00	1.9(-2)	$6.9(23)\pm2.4(23)$
G094.6028 6	21:39:58.05	+50:14:26.0	+28.2	0.29	65.8 ± 38.7	1.86	5.0(-3)	$1.8(23)\pm1.2(23)$
G094.6028 7	21:39:58.22	+50:14:21.7	+29.0	0.20	26.7±11.7	1.70	1.3(-2)	$4.5(23)\pm2.6(23)$
G094.6028 8	21:39:58.59	+50:14:20.5	+28.9	0.30	16.7 ± 4.4	1.32	1.8(-2)	$6.4(23)\pm 2.6(23)$
G100.38 1 C	22:16:10.37	+52:21:34.1	-37.4	0.13	68.2 ± 0.7	7.28	1.8(-2)	$6.5(23)\pm1.3(23)$
G100.38 2	22:16:10.48	+52:21:36.7	-37.3	0.16	37.4 ± 0.0	1.21	5.9(-3)	$2.1(23)\pm4.2(22)$
G100.38 3	22:16:10.42	+52:21:37.1	-37.4	0.18	42.3 ± 37.6	1.20	5.1(-3)	$1.8(23)\pm1.9(23)$
G100.38 4	22:16:10.58	+52:21:33.3	-37.6	0.18	14.7 ± 6.1	0.99	1.6(-2)	$5.5(23)\pm 3.4(23)$
G100.38 5	22:16:10.62	+52:21:36.1	-37.5	0.20	20.0 ± 10.0	0.67	7.0(-3)	$2.5(23)\pm1.7(23)$
G108.75 1 C	22:58:47.41	+58:45:02.0	-51.0	0.15	111.2 ± 8.2	16.01	1.8(-2)	$6.2(23)\pm1.3(23)$

Notes. $a(b) = a \times 10^{b}$.

Article number, page 30 of 81

Position	α	δ	$v_{\rm LSR}$	$\sigma_{ m line}$	$T_{\rm kin}$	$I_{1.37mm}$	τ_{v}^{cont}	$N(\mathrm{H}_2)$
	J(2000)	J(2000)	$({\rm km}{\rm s}^{-1})$	(K)	(K)	(mJy beam ⁻¹)		(cm^{-2})
G108.75.2 C	22:58:46.78	+58:45:03.6	-50.7	0.14	82.9+ 2.6	7.81	1.2(-2)	4.2(23)+8.4(22)
G108 75 3	22.58.46.97	+58.45.049	-50.9	0.16	39 1+17 5	6.24	21(-2)	7.6(23)+4.2(23)
G108.75 4	22:50:40.27	158.45.04.6	50.9	0.16	22.0 ± 5.0	0.24 2.14	1.1(-2)	$1.0(23)\pm1.2(23)$
G108.75 4	22.38.47.00	+ 50.45.04.0	-50.8	0.10	22.9 ± 3.9	2.14	1.4(-2)	$4.9(23)\pm1.0(23)$
G108.75 5	22:58:46.89	+58:45:04.3	-50.8	0.13	$155./\pm/5.4$	0.97	7.5(-4)	$2.7(22)\pm1.4(22)$
G108.75 6	22:58:47.38	+58:45:03.5	-50.7	0.14	28.8 ± 17.2	0.97	4.7(-3)	$1.7(23)\pm1.2(23)$
G138.2957 1	03:01:31.28	+60:29:12.7	-38.9	0.16	31.3 ± 6.7	6.28	3.0(-2)	$1.1(24)\pm 3.4(23)$
G138.2957 2	03:01:31.41	+60:29:12.4	-39.2	0.19	34.8 ± 16.4	3.78	1.6(-2)	$5.6(23) \pm 3.3(23)$
G138 2957 3	03.01.31 49	+60.29.12.2	-393	0 1 9	149 3+28 4	3 1 3	2.7(-3)	9.6(22)+2.7(22)
G138 2957 4	03:01:31 36	$\pm 60.29.185$	-38.0	0.20	10.3 ± 5.7	2.64	23(-2)	81(23)+35(23)
$C_{129} 2057 5$	02:01:21 40	+60.20.10.3	20.0	0.20	17.5 ± 5.7 25.1 ± 12.9	2.04	2.3(2)	$0.1(23)\pm 3.3(23)$ 2 2(22) $\pm 1.6(22)$
G138.2937 3	03:01:31.40	+00.29.19.3	-38.5	0.20	55.1 ± 12.8	2.28	9.4(-3)	$5.5(25)\pm1.0(25)$
G138.29576	03:01:31.28	+60:29:13.9	-38.0	0.16	/4.9±14.1	2.27	4.1(-3)	$1.4(23)\pm4.1(22)$
G138.29577	03:01:31.28	+60:29:10.6	-39.4	0.16	117.1±41.7	2.19	2.4(-3)	$8.7(22)\pm 3.7(22)$
G138.2957 8	03:01:32.16	+60:29:18.7	-38.6	0.29	41.1±11.1	2.02	7.0(-3)	$2.5(23)\pm9.1(22)$
G138.2957 9	03:01:31.46	+60:29:18.9	-37.3	0.21	63.6±14.3	1.87	4.0(-3)	$1.4(23)\pm 4.5(22)$
G138.2957 10	03:01:32.15	+60:29:19.9	-37.2	0.26	20.8 ± 4.9	1.85	1.4(-2)	$5.1(23) \pm 1.8(23)$
G138 2957 11	03.01.32.25	+60.29.177	-38.5	0.35	20.0+2.6	1.80	15(-2)	53(23)+14(23)
G138 2057 12	03:01:30.67	$\pm 60.29.17.0$	-36.9	0.22	20.0 ± 2.0 27 4 ± 14.6	1.00	80(-3)	$28(23)\pm10(23)$
C120.0001.1	02.07.24.40	+00.29.12.9	-30.9	0.22	27.4 ± 14.0	12.40	$\frac{6.0(-3)}{6.7(-2)}$	$\frac{2.0(23)\pm1.9(23)}{2.4(22)\pm7.4(22)}$
G139.9091 1	03:07:24.49	+38:30:42.8	-39.8	0.27	257.5±59.1	13.49	0.7(-3)	$2.4(23)\pm7.4(22)$
G139.9091 2	03:07:24.58	+58:30:53.0	-39.7	0.21	298.3 ± 0.0	6.63	2.8(-3)	$1.0(23)\pm2.0(22)$
G139.9091 3	03:07:23.94	+58:30:52.5	-39.8	0.24	57.2 ± 61.6	1.79	4.3(-3)	$1.5(23)\pm1.8(23)$
G139.9091 4	03:07:24.49	+58:30:50.3	-40.8	0.30	55.3 ± 43.1	1.59	4.0(-3)	$1.4(23)\pm1.2(23)$
G075.78 1 C	20:21:44.02	+37:26:37.5	- 6.9	0.38	176.8 ± 0.3	71.67	6.2(-2)	$2.1(24)\pm4.3(23)$
G075 78 2	20.21.43.95	+37.26.393	- 60	0.30	73 1+28 6	13.88	3.0(-2)	10(24)+49(23)
G075 78 3	20.21.13.95 20.21.44.10	$\pm 37.26.30.3$	- 67	0.30	617 ± 20.0	4 20	1.0(-2)	$3.6(23)\pm 1.6(23)$
C075.78.4	20.21.44.10	+27.26.29.0	- 0.7	0.51	04.7 ± 23.2	4.20	1.0(-2)	$5.0(25)\pm1.0(25)$
GU/5.78 4	20:21:44.28	+37:20:38.0	- /.1	0.30	2/0.8± 3.2	4.10	2.2(-3)	$7.8(22)\pm1.0(22)$
IRAS 21078 1 C	21:09:21.71	+52:22:37.1	- 6.5	0.40	135.5 ± 0.2	36.69	4.7(-2)	$1.6(24)\pm 3.3(23)$
IRAS 21078 2 C	21:09:21.79	+52:22:35.7	- 6.8	0.25	121.9 ± 3.4	22.95	3.3(-2)	$1.1(24)\pm 2.3(23)$
IRAS 21078 3	21:09:21.76	+52:22:36.8	- 5.4	0.29	120.5 ± 15.1	21.21	3.0(-2)	$1.1(24)\pm 2.6(23)$
IRAS 21078 4	21:09:22.12	+52:22:34.6	- 3.9	0.21	46.3±14.3	20.09	8.3(-2)	$2.8(24) \pm 1.1(24)$
IRAS 21078 5	21:09:21.85	+52:22:32.0	- 5.4	0.27	249.0 + 9.1	19.24	1.3(-2)	4.6(23)+9.3(22)
IRAS 21078 6	21.09.21.65	+52.22.386	- 63	0.25	61 8+ 3 8	19.06	5.6(-2)	20(24)+41(23)
IRAS 21078 7	21:09:21:05	$\pm 52.22.30.0$	- 46	0.25	206.0 ± 23.3	16.40	1.3(-2)	$4.8(23)\pm1.1(23)$
ID A C 21070 9	21.00.21.00	152.22.30.7	4 .0	0.25	200.7±25.5	12.09	1.3(2)	$+.0(23)\pm1.1(23)$ 1 1(24) + 5 6(22)
IRAS 21078 8	21:09:21.90	+32:22:33.4	- 3.2	0.20	77.7±33.0	15.98	5.2(-2)	$1.1(24)\pm 3.0(23)$
IRAS 21078 9	21:09:22.04	+52:22:35.2	- 5.5	0.26	229.5±18.5	10.08	7.4(-3)	$2.6(23)\pm 5.7(22)$
IRAS 21078 10	21:09:21.82	+52:22:38.0	- 6.0	0.25	115.7 ± 36.4	9.99	1.5(-2)	$5.3(23)\pm2.0(23)$
IRAS 21078 11	21:09:21.37	+52:22:38.1	- 6.0	0.32	229.1 ± 59.2	5.16	3.8(-3)	$1.3(23)\pm4.4(22)$
IRAS 21078 12	21:09:21.83	+52:22:34.0	- 5.3	0.28	41.4 ± 8.6	4.03	1.8(-2)	$6.4(23)\pm 2.0(23)$
IRAS 21078 13	21:09:21.83	+52:22:33.0	- 5.3	0.33	298.1 ± 0.9	3.96	2.2(-3)	$7.9(22) \pm 1.6(22)$
IRAS 21078 14	21.09.2149	+52.22.388	- 58	0.28	286 2+ 2.2	3 29	19(-3)	68(22)+14(22)
NGC7538 IPS9 1 C	23:14:01.75	$\pm 61.27.10.8$		0.20	200.5± 0.0	46.00	$\frac{1.7(-2)}{3.7(-2)}$	$13(24)\pm 26(23)$
NGC7528 IDS0 2	23.14.01.75	+61.27.19.0	-30.3	0.04	200.3 ± 0.9	40.09	5.7(-2)	$1.3(2+)\pm 2.0(23)$ $1.9(22)\pm 4.1(22)$
NGC / 538 IR 59 2	23:14:02.05	+01:27:19.7	-57.7	0.20	189.4 ± 18.4	0.12	5.2(-3)	$1.8(23)\pm4.1(22)$
NGC/5381RS9 3	23:14:02.21	+61:27:18.9	-58.4	0.27	194.6±21.8	4.65	3.8(-3)	$1.4(23)\pm 3.1(22)$
NGC7538 IRS9 4	23:14:02.15	+61:27:20.1	-57.9	0.27	104.2 ± 52.4	4.44	7.0(-3)	$2.5(23)\pm1.4(23)$
NGC7538 IRS9 5	23:14:01.64	+61:27:21.2	-58.5	0.22	263.4± 7.7	3.27	2.0(-3)	$7.0(22)\pm1.4(22)$
NGC7538 IRS9 6	23:14:01.89	+61:27:20.9	-58.8	0.32	84.5±22.5	2.81	5.5(-3)	$2.0(23)\pm6.8(22)$
NGC7538 IRS9 7	23:14:02.02	+61:27:20.6	-58.5	0.27	203.8 ± 29.4	2.71	2.1(-3)	$7.6(22) \pm 1.9(22)$
NGC7538 IRS9 8	23.14.02.34	+61.27.198	-58.0	0.19	134 8+70 6	2.33	28(-3)	1.0(23)+5.8(22)
\$87 IPS1 1 C	10:46:20.64	+24.35.38.8	+22.0	0.17	13.0 ± 70.0 112.0± 5.2	37.31	$\frac{2.8(-2)}{4.8(-2)}$	$1.0(23)\pm3.0(22)$ $1.7(24)\pm3.4(23)$
S07 IDS1 2	10.46.20.50	124.35.36.6	122.5	0.01	84.4 ± 27.0	8 24	+.0(-2)	$1.7(2+)\pm 3.4(23)$ $5.0(22)\pm 2.0(22)$
587 IRS1 2	19:40:20.39	+24:55:50.0	+22.3	0.58	64.4±27.0	8.24 6.77	1.4(-2)	$5.0(25)\pm 2.0(25)$
S87 IRS1 3	19:46:20.13	+24:35:29.4	+22.0	0.32	63.5 ± 40.1	6.//	1.6(-2)	$5.5(23)\pm4.0(23)$
S87 IRS1 4	19:46:19.87	+24:35:28.4	+22.2	0.28	28.4 ± 7.4	6.60	3.9(-2)	$1.3(24)\pm5.0(23)$
S87 IRS1 5	19:46:20.02	+24:35:28.8	+21.2	0.25	53.2 ± 9.9	5.92	1.7(-2)	$5.9(23) \pm 1.7(23)$
S87 IRS1 6	19:46:20.61	+24:35:37.4	+22.9	0.48	100.3 ± 52.1	5.42	7.7(-3)	$2.7(23) \pm 1.6(23)$
S87 IRS1 7	19:46:20.22	+24:35:32.0	+22.7	0.24	69.9 ± 16.4	3.31	6.9(-3)	$2.4(23) \pm 7.9(22)$
S87 IRS1 8	19:46:19 94	+24:35:30.2	+21.5	0.24	107.7+60.5	3.22	4.2(-3)	1.5(23)+9.3(22)
S87 IR S1 0	19.46.10.00	+24.35.30.2	+21.8	0.35	186.6 ± 43.0	2.58	1.2(-3)	$68(22) \pm 21(22)$
\$106.1	<u>17.70.17.77</u>	+27.00.477	1 21.0	0.33	61 2 42 4	150 47	$\frac{1.7(-5)}{5.2(-1)}$	$\frac{0.0(22)\pm2.1(22)}{1.4(25)\pm1.4(25)}$
S100 I	20.27.20.77	+37.22:47.7	+ 0.5	0.24	04.3 ± 03.4	130.47	J.2(-1)	$1.4(23)\pm1.0(23)$
51062	20:27:25.49	+57:22:48.4	- 0.5	0.57	124.8 ± 30.8	50.13	6.9(-2)	$2.4(24)\pm7.7(23)$
S106 3	20:27:25.51	+37:22:46.9	- 0.8	0.75	64.2±12.5	19.13	5.3(-2)	$1.8(24)\pm 5.3(23)$

Notes. $a(b) = a \times 10^{b}$.

Article number, page 31 of 81

Position	α	δ	v _{LSR}	$\sigma_{\rm line}$	T _{kin}	<i>I</i> _{1.37mm}	$ au_{v}^{\mathrm{cont}}$	<i>N</i> (H ₂)
	J(2000)	J(2000)	$({\rm km}{\rm s}^{-1})$	(K)	(K)	(mJy beam ⁻¹)		(cm^{-2})
S106 4	20:27:26.99	+37:22:50.5	- 2.4	0.25	96.7±38.8	8.05	1.4(-2)	5.0(23)±2.3(23)
W3 H2O 1	02:27:03.85	+61:52:25.0	-47.1	0.54	124.2 ± 28.3	456.29	1.3(0)	2.5(25)±7.9(24)
W3 H2O 2	02:27:03.90	+61:52:24.3	-46.8	0.41	81.0± 9.1	317.19	1.5(0)	$2.8(25)\pm6.5(24)$
W3 H2O 3 C	02:27:04.71	+61:52:24.7	-51.7	0.97	176.6±10.9	170.41	2.0(-1)	$6.6(24) \pm 1.4(24)$
W3 H2O 4 C	02:27:04.55	+61:52:24.7	-49.7	1.27	166.4 ± 9.0	161.32	2.1(-1)	$6.6(24) \pm 1.4(24)$
W3 H2O 5	02:27:04.42	+61:52:25.4	-48.1	0.42	109.0 ± 21.4	42.75	8.0(-2)	2.7(24)±7.8(23)
W3 IRS4 1 C	02:25:31.35	+62:06:20.8	-44.7	0.44	173.0 ± 15.4	51.50	5.6(-2)	$1.9(24)\pm4.3(23)$
W3 IRS4 2	02:25:31.97	+62:06:23.7	-46.6	0.63	48.2 ± 10.6	20.76	9.0(-2)	$3.1(24)\pm9.6(23)$
W3 IRS4 3	02:25:31.50	+62:06:22.2	-43.9	0.46	61.3±19.4	16.36	5.3(-2)	$1.9(24) \pm 7.4(23)$
W3 IRS4 4	02:25:31.04	+62:06:19.8	-44.6	0.45	251.5± 9.5	10.46	7.6(-3)	$2.7(23)\pm5.5(22)$
W3 IRS4 5	02:25:31.50	+62:06:22.9	-44.5	0.43	27.9 ± 3.1	10.02	8.1(-2)	$2.8(24)\pm6.7(23)$
W3 IRS4 6	02:25:31.62	+62:06:24.9	-44.6	0.39	84.8 ± 29.0	5.91	1.3(-2)	$4.7(23)\pm2.0(23)$

Notes. $a(b) = a \times 10^{b}$.

Appendix B: Spectral line properties

The line properties of the detected spectral lines analyzed in this study are summarized in Table B.1.

Table B.1: Properties of the analyzed spectral lines taken from Splatalogue (Remijan et al. 2007). The entries are taken from the CDMS and JPL catalogs (Müller et al. 2005; Pickett et al. 1998). Blended lines are indicated by a ‡. The mean and maximum line optical depth for each transition fitted with XCLASS are computed considering all 120 positions (see Sect. 4.2).

			Einstein	Upper		Li	ne
		Frequency	coefficient	Energy Level		Optical	l Depth
Molecule	Ouantum Numbers	ν	$\log(A_{\rm ul})$	$E_{\rm u}/k_{\rm B}$	Catalog	$\tau_{\rm magn}^{\rm line}$	$\tau_{\rm max}^{\rm line}$
	C C	(GHz)	$(\log s^{-1})$	(K)	U	mean	шах
¹³ CO	2 - 1	220.399	-6.22	15.9	JPL	1.5(0)	2.1(2)
C ¹⁸ O	2 - 1	219 560	-6.22	15.8	IPL	$\frac{11(-1)}{11(-1)}$	$\frac{54(0)}{54(0)}$
$\frac{0}{50}$	$\frac{5}{65-54}$	219.900	-3.87	35.0	IPL	$\frac{1.1(-1)}{2.7(-1)}$	$\frac{3.1(0)}{8.3(0)}$
005	$\frac{0.5}{18-17}$	219.913	-4.52	99.8	IPI	$\frac{2.7(-1)}{4.3(-1)}$	$\frac{0.3(0)}{2.4(1)}$
$\frac{000}{0^{13}}$	$\frac{10}{18} - 17$	218.100	-4.52	00.5	IDI	$\frac{-4.3(-1)}{-2.0(-3)}$	$\frac{2.4(1)}{3.4(-1)}$
<u> </u>	$\frac{10-17}{22715-23}$	210.199	-4.67	352.8	JI L IPI	$\frac{2.0(-3)}{2.8(-2)}$	$\frac{5.4(-1)}{5.6(-1)}$
<u>34</u> SO	$\frac{227,15 - 236,18}{11}$	219.270	2.06	60.2		$\frac{2.0(-2)}{1.1(-2)}$	$\frac{J.0(-1)}{1.1(-1)}$
$\frac{30_2}{DCN}$	$\frac{11_{1,11} - 10_{0,10}}{2 2}$	219.333	$\frac{-3.90}{3.34}$	20.0	CDMS	$\frac{1.1(-2)}{0.7(-2)}$	$\frac{1.1(-1)}{2.2(1)}$
	$\frac{3-2}{3-2}$	217.239	-3.54	20.9		$\frac{9.7(-2)}{6.2(-1)}$	$\frac{2.2(1)}{2.5(1)}$
	$3_{0,3} - 2_{0,2}$	210.222	-3.33	21.0 69.1	JL IDI	0.2(-1)	2.3(1) 5.8(0)
	$3_{2,2} - 2_{2,1}$	218.470	-3.80	68 1	JFL IDI	1.3(-1) 1.2(1)	3.6(0)
$\frac{\Pi_2 CO}{U^{13} CO}$	$\frac{3_{2,1}-2_{2,0}}{2}$	210.700	-3.80	22.0		$\frac{1.3(-1)}{1.5(-2)}$	$\frac{0.3(0)}{7.0(-1)}$
$\frac{H_2^{-1}CO}{UNCO}$	$\frac{3_{1,2}-2_{1,1}}{10}$	219.909	-3.39	32.9	JPL CDMC	$\frac{1.3(-2)}{(-2)}$	$\frac{7.9(-1)}{2.2(0)}$
HNCO	$10_{1,10} - 9_{1,9}$	218.981	-3.85	101.1	CDMS	0.3(-2)	2.3(0)
HNCO	$10_{3,8} - 9_{3,7}$	219.057	-3.92	433.0	CDMS	1.0(-2)	1.7(-1)
HNCO	$10_{3,7} - 9_{3,6}$	219.037	-3.92	455.0	CDMS	1.0(-2)	1./(-1)
HNCO	$10_{2,9} - 9_{2,8}$	219.734	-3.87	228.3	CDMS	5.8(-2)	9.9(-1)
HNCO	$10_{2,8} - 9_{2,7}$	219.757	-3.87	220.5	CDMS	0.3(-2)	9.9(-1)
HNCO	$10_{0,10} - 9_{0,9}$	219.798	-3.83	58.0 101.5	CDMS	1.3(-1)	3.9(0)
	$10_{1,9} - 9_{1,8}$	220.383	-3.84	101.5		9.1(-2)	$\frac{2.3(0)}{1.2(0)}$
$\frac{\Pi C_3 N}{\Pi C_3 N}$	24 - 25	218.525	-3.08	151.0	JPL	8.3(-2)	1.5(0)
$HC_3N; v_7 = 1^*$	24 - 23, l = 1e	218.861	-3.08	452.1	CDMS	-	-
$HC_3N; v_7 = 1$	24 - 23, l = 1f	219.174	-3.08	452.3	CDMS	$\frac{2.5(-2)}{2.4(-2)}$	$\frac{5.8(-1)}{(-1)}$
$HC_3N; v_7 = 2$	24 - 23, l = 0	219.675	-3.08	773.5	CDMS	3.4(-3)	6.8(-2)
$HC_3N; v_7 = 2$	24 - 23, l = 2e	219.707	-3.08	//6.8	CDMS	4.2(-3)	6.6(-2)
$HC_3N; v_7 = 2^*$	24 - 23, l = 2f	219.742	-3.08	776.8	CDMS	-	-
HCC ¹⁵ CN	24 - 23	217.420	-3.09	130.4	CDMS	1.1(-3)	$\frac{1.8(-2)}{2}$
CH ₃ OH	$20_{1,19} - 20_{0,20}E$	217.887	-4.47	508.4	CDMS	1.4(-2)	5.3(-1)
CH ₃ OH	$4_{2,3} - 3_{1,2}E$	218.440	-4.33	45.5	CDMS	5.7(-1)	4.0(1)
CH ₃ OH	$25_{3,23} - 24_{4,20}E$	219.984	-4.69	802.2	CDMS	1.5(-3)	9.6(-2)
CH ₃ OH	$23_{5,18} - 22_{6,17}E$	219.994	-4.76	775.9	CDMS	1.5(-3)	9.4(-2)
CH ₃ OH	$8_{0,8} - 7_{1,6}E$	220.079	-4.60	96.6	CDMS	9.5(-2)	5.0(0)
CH ₃ OH ⁺	$10_{5,6} - 11_{4,8}E$	220.401	-4.95	251.6	CDMS	-	-
$CH_3OH; v_t = 1$	$6_{1,5} - 7_{2,5}A$	217.299	-4.37	373.9	CDMS	4.1(-2)	1.6(0)
$CH_3OH; v_t = 1$	$15_{6,9} - 16_{5,11}A$	217.643	-4.72	745.6	CDMS	1.0(-2)	1.5(-1)
$CH_3OH; v_t = 1$	$15_{6,10} - 16_{5,12}A$	217.643	-4.72	/45.6	CDMS	1.0(-2)	$\frac{1.5(-1)}{1.5(-1)}$
CH ₃ CN	$12_8 - 11_8$	220.476	-3.45	525.6	JPL	7.4(-3)	4.0(-1)
CH ₃ CN	$12_7 - 11_7$	220.539	-3.38	418.6	JPL	2.2(-2)	7.7(-1)
CH ₃ CN	$12_6 - 11_6$	220.594	-3.32	325.9	JPL	9.2(-2)	2.7(0)
CH ₃ CN	$12_5 - 11_5$	220.641	-3.28	247.4	JPL	5.9(-2)	2.1(0)
CH_3CN	$12_4 - 11_4$	220.679	-3.25	183.1	JPL	1.1(-1)	2.7(0)
CH_3CN	$12_3 - 11_3$	220.709	-3.22	133.2	JPL	2.7(-1)	8.2(0)
CH_3CN	$12_2 - 11_2$	220.730	-3.21	97.4	JPL	2.1(-1)	4.7(0)
CH ₃ CN	$12_1 - 11_1$	220.743	-3.20	/6.0	JPL	4.5(-1)	6.7(0)
CH ₃ CN	$12_0 - 11_0$	220.747	-3.20	08.9	JPL	$\frac{3.3(-1)}{0.9(-1)}$	$\frac{0.7(0)}{2.1(-2)}$
$CH_3^{-1}CN$	$12_6 - 11_6$	220.486	-3.16	325.9	JPL	9.8(-4)	3.1(-2)
$CH_3^{13}CN$	$12_5 - 11_5$	220.532	-3.12	247.4	JPL	5.7(-3)	4.8(-1)
$CH_{3}^{13}CN$	$12_4 - 11_4$	220.570	-3.09	183.1	JPL	1.3(-3)	3.9(-2)
$CH_{3}^{13}CN$	$12_3 - 11_3$	220.600	-3.06	133.1	JPL	1.5(-2)	6.4(-1)
$CH_3^{13}CN$	$12_2 - 11_2$	220.621	-3.05	97.4	JPL	2.0(-3)	6.3(-2)
CH ₃ ¹³ CN	$12_1 - 11_1$	220.634	-3.04	76.0	JPL	1.4(-2)	1.0(0)
CH ¹³ CN	$12_0 - 11_0$	220.638	-3.03	68.8	JPL	6.4(-2)	2.1(0)

Notes. $a(b) = a \times 10^{b}$.

Appendix C: Temperature maps

Figure C.1 shows the H_2CO and CH_3CN temperature maps derived with XCLASS for each region (Sect. 3.2). Each core position is marked in red and the derived outer radii from the radial temperature profiles are indicated by dashed red circles.



Fig. C.1: Temperature maps derived with XCLASS. Each panel shows in color the temperature map (left: H₂CO, right: CH₃CN) and in black contours the 1.37 mm continuum emission. The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with steps increasing by a factor of 2 (see Table 2 for values of σ_{cont} for each region). Each core is marked in red and the dashed red circle indicates the outer radius of the radial temperature fit (Sect. 3.2). The beam size is shown in the bottom left corner in each panel. The pink bar in the top left corner indicates a linear spatial scale of 10 000 au. The primary beam size is indicated by a black circle and for regions with no extended H₂CO temperature map a smaller field of view is shown.



Fig. C.1 (cont.): Temperature maps derived with XCLASS. Each panel shows in color the temperature map (left: H₂CO, right: CH₃CN) and in black contours the 1.37 mm continuum emission. The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with steps increasing by a factor of 2 (see Table 2 for values of σ_{cont} for each region). Each core is marked in red and the dashed red circle indicates the outer radius of the radial temperature fit (Sect. 3.2). The beam size is shown in the bottom left corner in each panel. The pink bar in the top left corner indicates a linear spatial scale of 10 000 au. The primary beam size is indicated by a black circle and for regions with no extended H₂CO temperature map a smaller field of view is shown.


Fig. C.1 (cont.): Temperature maps derived with XCLASS. Each panel shows in color the temperature map (left: H₂CO, right: CH₃CN) and in black contours the 1.37 mm continuum emission. The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with steps increasing by a factor of 2 (see Table 2 for values of σ_{cont} for each region). Each core is marked in red and the dashed red circle indicates the outer radius of the radial temperature fit (Sect. 3.2). The beam size is shown in the bottom left corner in each panel. The pink bar in the top left corner indicates a linear spatial scale of 10 000 au. The primary beam size is indicated by a black circle and for regions with no extended H₂CO temperature map a smaller field of view is shown.



Fig. C.1 (cont.): Temperature maps derived with XCLASS. Each panel shows in color the temperature map (left: H₂CO, right: CH₃CN) and in black contours the 1.37 mm continuum emission. The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with steps increasing by a factor of 2 (see Table 2 for values of σ_{cont} for each region). Each core is marked in red and the dashed red circle indicates the outer radius of the radial temperature fit (Sect. 3.2). The beam size is shown in the bottom left corner in each panel. The pink bar in the top left corner indicates a linear spatial scale of 10 000 au. The primary beam size is indicated by a black circle and for regions with no extended H₂CO temperature map a smaller field of view is shown.



Fig. C.1 (cont.): Temperature maps derived with XCLASS. Each panel shows in color the temperature map (left: H₂CO, right: CH₃CN) and in black contours the 1.37 mm continuum emission. The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with steps increasing by a factor of 2 (see Table 2 for values of σ_{cont} for each region). Each core is marked in red and the dashed red circle indicates the outer radius of the radial temperature fit (Sect. 3.2). The beam size is shown in the bottom left corner in each panel. The pink bar in the top left corner indicates a linear spatial scale of 10 000 au. The primary beam size is indicated by a black circle and for regions with no extended H₂CO temperature map a smaller field of view is shown.



Fig. C.1 (cont.): Temperature maps derived with XCLASS. Each panel shows in color the temperature map (left: H₂CO, right: CH₃CN) and in black contours the 1.37 mm continuum emission. The dashed black contours show the $-5\sigma_{cont}$ emission and the solid black contours start at $5\sigma_{cont}$ with steps increasing by a factor of 2 (see Table 2 for values of σ_{cont} for each region). Each core is marked in red and the dashed red circle indicates the outer radius of the radial temperature fit (Sect. 3.2). The beam size is shown in the bottom left corner in each panel. The pink bar in the top left corner indicates a linear spatial scale of 10 000 au. The primary beam size is indicated by a black circle and for regions with no extended H₂CO temperature map a smaller field of view is shown.

Appendix D: Abundance histograms

Figure D.1 shows a histogram of the abundance ratio relative to $N(C^{18}O)$ for each molecule complementary to the column density histograms discussed in Sect. 4.2.



Fig. D.1: Abundance ratio histograms. Abundance ratio histograms of all 120 positions are shown in green (upper limits are not included). Separate abundance ratio histograms of the core and non core positions are indicated by the dash-dotted red and dashed blue lines, respectively.

Appendix E: Column density results

Table E.1, E.2, and E.3 show the derived column densities for all molecules fitted with XCLASS (Sect. 4.2). The upper and lower errors are estimated using the MCMC error estimation algorithm in XCLASS. If the fitting results do not match our defined criteria for a good fit (further explained in Sect. 4.2), the results are considered as upper limits.

Table E.1: Molecular column densities derived with XCLASS.

Position	¹³ CO	C ¹⁸ O	SO	OCS	SO ₂
IRAS 23033 1	$8.4(17)^{+9.1(16)}_{-1.2(17)}$	$5.3(16)^{+5.8(16)}_{-1.7(16)}$	$1.7(15)^{+1.7(15)}_{-3.5(14)}$	<9.5(14)	<1.1(16)
IRAS 23033 2	$7.6(17)^{+1.8(17)}_{-8.4(16)}$	$1.9(17)^{+1.0(17)}_{-3.9(16)}$	$2.5(16)^{+1.1(16)}_{-4.6(15)}$	$9.5(15)^{+5.0(15)}_{-1.3(15)}$	$3.4(16)^{+3.1(16)}_{-1.4(16)}$
IRAS 23033 3	$1.2(18)^{+1.7(17)}_{-1.0(17)}$	$2.6(17)^{+5.4(16)}_{-4.5(16)}$	$2.8(16)^{+1.6(15)}_{-1.5(15)}$	$2.4(16)^{+4.3(15)}_{-3.7(15)}$	$4.7(16)^{+3.2(16)}_{-1.4(16)}$
IRAS 23033 4	$8.2(17)^{+2.8(17)}_{-9.6(16)}$	$2.4(17)^{+6.4(16)}_{-6.0(16)}$	$2.3(15)^{+2.1(15)}_{-8.0(14)}$	<1.8(14)	<1.4(16)
IRAS 23033 5	$8.0(17)^{+2.0(17)}_{-7.4(16)}$	$1.3(17)^{+1.8(17)}_{-3.2(16)}$	$6.3(14)^{+1.2(14)}_{-5.0(14)}$	<3.9(14)	<1.3(16)
IRAS 23151 1	$1.8(18)^{+2.5(17)}_{-1.5(17)}$	$6.9(17)^{+1.0(17)}_{-7.4(16)}$	$4.0(16)^{+1.5(15)}_{-5.5(15)}$	$1.5(16)^{+3.3(15)}_{-1.5(15)}$	$5.1(17)^{+3.8(16)}_{-1.9(16)}$
IRAS 23151 2	$6.4(17)^{+6.9(16)}_{-1.1(17)}$	$2.1(17)^{+7.5(16)}_{-3.2(16)}$	$7.9(15)^{+1.8(15)}_{-1.2(15)}$	$7.6(14)^{+8.2(13)}_{-6.5(14)}$	<1.8(16)
IRAS 23151 3	$6.7(17)^{+1.2(17)}_{-1.0(17)}$	$4.0(17)^{+5.9(16)}_{-6.2(16)}$	$8.3(15)^{+5.2(15)}_{-1.9(15)}$	$2.0(15)^{+1.1(16)}_{-7.5(14)}$	<1.5(16)
IRAS 23151 4	$2.8(17)^{+1.1(17)}_{-5.6(16)}$	$2.3(17)^{+9.5(16)}_{-3.2(16)}$	$3.3(15)^{+1.1(15)}_{-3.8(14)}$	$7.1(14)^{+1.5(14)}_{-5.8(14)}$	<2.7(16)
IRAS 23151 5	$9.5(18)_{-8.9(16)}^{+4.8(16)}$	$1.3(17)^{+5.1(16)}_{-4.4(16)}$	$4.3(15)_{-9.4(14)}^{+1.2(15)}$	<3.7(14)	<1.5(16)
IRAS 23385 1	$9.3(17)^{+2.3(17)}_{-1.5(17)}$	$2.6(17)^{+1.4(17)}_{-3.5(16)}$	$1.1(16)^{+3.8(15)}_{-1.6(15)}$	$1.2(16)^{+1.7(15)}_{-2.1(15)}$	<1.5(16)
IRAS 23385 2	$6.2(17)^{+2.2(17)}_{-1.1(17)}$	$2.1(17)^{+1.6(17)}_{-2.8(16)}$	$2.2(15)^{+1.9(15)}_{-7.5(14)}$	$1.3(15)^{+6.2(15)}_{-6.9(14)}$	<1.7(18)
IRAS 23385 3	$4.3(17)^{+5.9(16)}_{-1.1(17)}$	$2.3(17)^{+1.5(17)}_{-3.5(16)}$	$5.6(15)^{+2.1(15)}_{-2.4(15)}$	$9.6(14)^{+1.9(15)}_{-4.2(14)}$	<2.0(16)
IRAS 23385 4	$1.0(18)^{+1.9(17)}_{-1.9(17)}$	$9.5(16)^{+1.2(17)}_{-2.8(16)}$	$4.6(15)^{+1.8(15)}_{-1.3(15)}$	$8.9(14)^{+1.8(14)}_{-7.2(14)}$	<1.6(16)
IRAS 23385 5	$5.8(17)^{+2.1(17)}_{-1.6(17)}$	$7.1(16)^{+8.2(16)}_{-3.3(16)}$	$4.3(15)^{+1.0(15)}_{-1.4(15)}$	<6.1(14)	<1.2(16)
IRAS 23385 6	$6.5(17)^{+1.5(17)}_{-1.7(17)}$	$1.2(17)^{+8.6(16)}_{-4.6(16)}$	$3.5(15)^{+1.7(15)}_{-8.0(14)}$	<2.6(14)	<1.1(16)
IRAS 23385 7	$1.3(18)^{+3.3(17)}_{-3.7(17)}$	$1.2(17)^{+7.4(16)}_{-5.3(16)}$	$1.4(16)^{+6.9(15)}_{-7.4(15)}$	<8.8(14)	<1.0(16)
IRAS 23385 8	$3.8(18)^{+5.4(17)}_{-6.8(17)}$	$8.4(16)^{+5.0(16)}_{-4.8(16)}$	$1.3(15)^{+4.9(14)}_{-3.6(14)}$	$1.6(15)^{+1.2(16)}_{-6.7(14)}$	<3.8(16)
AFGL 2591 1	$4.6(18)^{+3.1(17)}_{-2.4(17)}$	$4.1(17)^{+6.0(16)}_{-3.6(16)}$	$1.1(17)^{+3.8(15)}_{-6.8(15)}$	$4.1(16)^{+5.7(15)}_{-4.2(15)}$	$1.3(18)^{+2.2(17)}_{-1.3(17)}$
AFGL 2591 2	$6.6(17)^{+1.5(17)}_{-6.8(16)}$	$6.0(17)^{+9.7(16)}_{-5.9(16)}$	$1.3(15)^{+7.1(14)}_{-6.1(14)}$	<1.6(15)	<1.3(16)
AFGL 2591 3	$6.3(17)^{+2.0(17)}_{-1.0(17)}$	$2.7(17)^{+3.1(16)}_{-4.9(16)}$	$4.8(15)^{+2.0(16)}_{-2.8(15)}$	<1.7(17)	<2.4(16)
AFGL 2591 4	$9.8(17)_{-9.9(16)}^{+1.7(17)}$	$1.9(17)^{+8.3(16)}_{-2.2(16)}$	$6.3(15)^{+1.6(15)}_{-1.2(15)}$	<1.5(15)	<2.1(16)
CepA HW2 1	$1.1(18)^{+4.8(17)}_{-2.4(17)}$	$5.6(17)^{+1.4(17)}_{-6.2(16)}$	$1.4(17)^{+2.3(16)}_{-1.9(16)}$	$3.6(16)^{+9.3(15)}_{-2.0(15)}$	$1.3(18)^{+1.2(17)}_{-1.1(17)}$
CepA HW2 2	$1.1(18)^{+1.4(17)}_{-1.5(17)}$	$5.6(17)^{+4.8(16)}_{-9.9(16)}$	$5.1(16)^{+2.1(16)}_{-5.5(15)}$	$1.2(16)^{+2.5(16)}_{-2.1(15)}$	$3.3(17)^{+1.2(17)}_{-5.2(16)}$
CepA HW2 3	$2.2(18)^{+2.6(17)}_{-4.1(17)}$	$7.5(17)^{+5.5(16)}_{-5.0(16)}$	$3.0(16)^{+0.8(15)}_{-4.6(15)}$	$2.0(16)^{+5.5(15)}_{-2.5(15)}$	$9.1(16)^{+1.5(16)}_{-1.3(16)}$
CepA HW2 4	$9.9(18)^{+2.3(16)}_{-2.3(16)}$	$1.2(17)^{+5.3(16)}_{-5.9(16)}$	<1.1(14)	<1.0(14)	<1.7(16)
CepA HW2 5	$9.9(17)^{+1.3(17)}_{-1.1(17)}$	$2.9(17)^{+1.1(17)}_{-4.6(16)}$	$1.2(16)^{+1.4(15)}_{-2.7(15)}$	<7.5(14)	<1.0(16)
G084.9505 1	$5.9(17)^{+2.3(17)}_{-1.2(17)}$	$1.1(17)^{+9.3(16)}_{-3.1(16)}$	$2.7(15)^{+2.9(14)}_{-7.2(14)}$	$3.4(15)^{+2.0(15)}_{-7.4(14)}$	<8.2(17)
G084.9505 2	$2.5(17)^{+1.5(17)}_{-1.1(16)}_{+1.2(17)}$	$4.1(16)^{+1.4(16)}_{-2.9(16)}_{+4.7(16)}$	$1.0(15)^{+3.9(14)}_{-3.2(14)}$	<4.7(14)	<1.2(16)
G084.9505 3	$1.1(17)^{+1.2(17)}_{-1.4(16)}$	$5.1(16)^{+4.7(16)}_{-2.5(16)}_{+2.5(16)}$	$1.4(15)^{+1.2(15)}_{-4.5(14)}_{+6.7(14)}$	<1.5(14)	<1.0(16)
G084.9505 4	$1.5(17)^{+4.5(10)}_{-4.7(16)}$	$2.6(16)^{+2.5(10)}_{-4.1(15)}$	$1.6(15)^{+6.7(14)}_{-7.6(14)}$	<1.9(14)	<3.5(18)
G084.9505 5	$6.8(17)^{+4.1(17)}_{-1.5(17)}$	$9.3(16)^{+1.0(17)}_{-2.7(16)}$	$1.8(15)^{+0.2(14)}_{-1.2(15)}$	<2.3(14)	<1.2(16)
G084.9505 6	$2.2(17)^{+1.7(17)}_{-2.1(16)}_{+4.0(17)}$	$5.3(16)^{+2.4(10)}_{-2.7(16)}$	$2.0(15)^{+1.5(15)}_{-1.2(15)}$	$8.9(14)^{+3.0(15)}_{-4.2(14)}$	<4.0(16)
G084.9505 7	$6.7(17)^{+4.0(17)}_{-1.8(17)}$	$1.1(17)^{+7.4(10)}_{-5.2(16)}$	$2.0(15)^{+2.5(15)}_{-9.6(14)}$	$8.3(14)^{+1.7(13)}_{-4.9(14)}$	<1.0(16)
G084.9505 8	$4.1(17)^{+1.4(17)}_{-1.2(17)}$	$4.9(16)^{+4.5(16)}_{-2.3(16)}$	$4.2(14)^{+1.7(14)}_{-3.1(14)}$	<5.5(14)	<3.0(16)
G094.6028 1	$1.5(18)^{+3.5(17)}_{-1.0(17)}$	$3.1(16)^{+1.8(16)}_{-9.5(15)}$	$2.7(16)^{+4.0(15)}_{-3.5(15)}$	$3.2(16)^{+8.4(15)}_{-6.6(15)}$	$6.5(16)^{+2.9(16)}_{-2.8(16)}$
G094.6028 2	$3.4(17)^{+1.5(17)}_{-6.0(16)}$	$6.4(16)^{+1.9(16)}_{-4.1(16)}$	$1.0(15)^{+1.0(14)}_{-8.4(14)}$	<2.9(14)	<1.3(16)
G094.6028 3	$1.1(18)^{+2.3(17)}_{-4.9(17)}$	$1.2(17)^{+8.0(16)}_{-4.1(16)}$	$4.8(14)^{+1.2(15)}_{-1.6(14)}_{+4.2(14)}$	<1.3(14)	<1.3(16)
G094.6028 4	$4.0(17)^{+1.5(17)}_{-6.2(16)}$	$6.2(16)^{+4.0(16)}_{-2.9(16)}$	$1.3(15)^{+4.0(14)}_{-6.7(14)}$	<1.3(15)	<1.2(16)
G094.6028 5	$4.7(17)^{+3.0(16)}_{-1.2(17)}$	$5.3(16)^{+0.4(16)}_{-2.5(16)}$	$2.8(14)^{+4.1(14)}_{-4.1(13)}$	<4.0(14)	<1.0(16)
G094.6028 6	$1.3(18)^{+1.4(17)}_{-2.8(17)}$	$8.2(16)^{+3.8(16)}_{-2.7(16)}$	$1.0(15)^{+5.9(14)}_{-3.8(14)}$	<1.1(15)	<1.3(16)
G094.6028 7	$5.3(17)^{+1.1(17)}_{-9.4(16)}$	$5.7(16)^{+6.7(15)}_{-4.0(16)}$	$5.3(14)^{+4.2(13)}_{-4.1(14)}$	$8.8(14)^{+2.4(15)}_{-4.5(14)}$	<1.2(16)

Notes. $a(b) = a \times 10^{b}$.

Article number, page 43 of 81

Table E.1: d	continued.
--------------	------------

Position	¹³ CO	C ¹⁸ O	SO	OCS	SO ₂
G094.6028 8	$4.8(17)^{+3.3(17)}_{-7.7(16)}$	$4.8(16)^{+4.4(16)}_{-2.0(16)}$	<2.5(14)	<1.9(14)	<1.8(16)
G100.38 1	$6.6(17)^{+1.5(17)}_{-1.2(17)}$	$1.7(17)^{+1.6(17)}_{-5.0(16)}$	$1.6(16)^{+2.7(15)}_{-2.4(15)}$	$2.0(15)^{+4.0(15)}_{-1.5(15)}$	$4.3(16)^{+1.9(16)}_{-1.1(16)}$
G100.38 2	$5.5(17)^{+1.9(17)}_{-1.0(17)}$	$2.2(17)^{+1.7(17)}_{-6.9(16)}$	$1.8(15)^{+2.5(15)}_{-6.7(14)}$	<1.0(18)	<1.3(16)
G100.38 3	$3.6(17)^{+5.6(16)}_{-3.3(16)}$	$1.4(17)^{+7.8(16)}_{-3.5(16)}$	$8.4(14)^{+5.3(14)}_{-6.4(14)}$	<6.7(16)	<3.0(16)
G100.38 4	$7.1(16)^{+4.5(16)}_{-3.7(16)}$	$1.7(17)^{+6.8(16)}_{-6.7(16)}$	$2.6(15)^{+4.4(15)}_{-1.8(15)}$	<1.4(14)	<8.9(17)
G100.38 5	$2.7(17)^{+3.0(17)}_{-1.6(17)}$	$9.0(16)^{+7.7(16)}_{-4.8(16)}$	$1.1(15)^{+1.1(14)}_{-9.2(14)}$	<4.4(14)	<1.2(16)
G108.75 1	$6.9(17)^{+3.1(16)}_{-8.5(16)}$	$2.1(17)^{+1.8(17)}_{-2.9(16)}$	$3.8(15)^{+2.5(14)}_{-6.7(14)}$	$8.9(14)^{+4.6(14)}_{-7.2(14)}$	<1.9(16)
G108.75 2	$4.4(17)^{+1.6(17)}_{-2.3(17)}$	$5.2(16)^{+1.4(16)}_{-3.2(16)}$	$1.0(15)^{+6.5(14)}_{-4.6(14)}$	<1.9(14)	<2.7(18)
G108.75 3	$4.3(17)^{+3.8(16)}_{-6.3(16)}$	$5.8(16)^{+4.6(16)}_{-3.0(16)}$	$6.0(14)^{+6.6(14)}_{-3.1(14)}$	<1.4(14)	<1.1(16)
G108.75 4	$4.5(17)^{+7.8(16)}_{-4.5(16)}$	$1.3(17)^{+1.5(17)}_{-6.1(16)}$	$5.2(14)^{+7.3(14)}_{-2.3(14)}$	<5.4(15)	<1.0(16)
G108.75 5	$4.8(17)^{+5.6(17)}_{-1.4(17)}$	$5.0(16)^{+4.9(16)}_{-2.5(16)}$	$1.2(15)^{+3.1(15)}_{-6.9(14)}$	$3.5(14)^{+9.8(14)}_{-7.3(13)}$	<1.0(16)
G108.75 6	$8.8(17)^{+2.7(17)}_{-1.2(17)}$	$1.0(17)^{+1.1(17)}_{-3.9(16)}$	$1.2(15)^{+2.8(15)}_{-6.7(14)}$	<2.6(14)	<1.1(16)
G138.2957 1	$6.3(17)^{+1.9(17)}_{-6.7(16)}$	$6.1(16)^{+4.1(16)}_{-2.6(16)}$	$1.1(15)^{+1.4(14)}_{-9.8(14)}$	<2.1(14)	<1.2(16)
G138.2957 2	$4.0(17)^{+5.9(16)}_{-2.9(16)}$	$1.5(17)^{+7.0(16)}_{-4.0(16)}$	$1.0(15)^{+1.2(15)}_{-8.1(14)}$	<9.9(17)	<1.0(16)
G138.2957 3	$7.4(17)^{+1.8(17)}_{-1.7(17)}$	$1.7(17)^{+1.1(17)}_{-5.7(16)}$	$1.4(15)^{+1.6(15)}_{-6.9(14)}$	<5.5(14)	<2.4(18)
G138.2957 4	$1.3(18)^{+7.0(17)}_{-4.5(17)}$	$1.9(17)^{+1.0(17)}_{-8.0(16)}$	$1.8(15)^{+2.6(15)}_{-6.8(14)}$	<5.0(14)	<1.5(16)
G138.2957 5	$6.0(17)^{+1.9(17)}_{-8.2(16)}$	$1.7(17)^{+9.6(16)}_{-6.1(16)}$	$9.9(14)^{+8.5(14)}_{-6.0(14)}$	<1.0(14)	<2.4(16)
G138.2957 6	$6.4(17)^{+3.2(17)}_{-1.2(17)}$	$4.2(16)^{+5.9(16)}_{-1.3(16)}$	$8.4(14)^{+2.8(15)}_{-3.9(14)}$	<4.1(14)	<6.3(18)
G138.2957 7	$6.2(17)^{+1.7(17)}_{-1.6(17)}$	$6.1(16)^{+3.9(16)}_{-1.2(16)}$	$4.8(14)^{+5.6(14)}_{-2.9(14)}$	<1.7(14)	<1.1(16)
G138.2957 8	$7.2(17)^{+1.2(17)}_{-7.5(16)}$	$1.0(17)^{+8.8(16)}_{-4.7(16)}$	$1.2(15)^{+9.7(14)}_{-4.5(14)}$	<6.1(14)	<2.0(16)
G138.2957 9	$7.9(17)^{+1.2(17)}_{-1.8(17)}$	$1.9(17)^{+1.2(17)}_{-6.3(16)}$	$7.6(14)^{+8.4(14)}_{-1.7(14)}$	<8.7(14)	$1.5(16)^{+9.8(15)}_{-1.3(15)}$
G138.2957 10	$5.5(17)^{+0.4(17)}_{-4.6(16)}$	$7.1(16)^{+4.8(16)}_{-3.8(16)}$	$1.1(15)^{+8.5(14)}_{-6.2(14)}$	<7.2(14)	<1.7(16)
G138.2957 11	$1.1(18)^{+2.6(17)}_{-9.6(16)}$	$1.1(17)^{+2.0(17)}_{-3.9(16)}$	$9.2(14)^{+9.7(14)}_{-4.7(14)}$	<9.2(15)	<8.9(18)
G138.2957 12	$3.9(17)^{+2.7(17)}_{-1.2(17)}$	$1.1(17)^{+7.9(16)}_{-4.0(16)}$	$1.3(15)^{+0.2(14)}_{-1.0(15)}$	<3.2(14)	<1.7(18)
G139.9091 1	$3.4(17)^{+1.4(17)}_{-5.6(16)}_{+4.0(17)}$	$2.1(16)^{+1.4(10)}_{-3.4(15)}_{+4.6(16)}$	<3.8(14)	<1.1(14)	<8.3(18)
G139.9091 2	$1.4(18)^{+4.0(17)}_{-2.6(17)}$	$5.3(16)^{+4.0(10)}_{-2.1(16)}$	$1.9(15)^{+1.2(15)}_{-5.2(14)}$	<3.0(14)	<1.6(16)
G139.9091 3	$6.6(17)^{+9.5(10)}_{-7.0(16)}$	$5.6(16)^{+5.1(10)}_{-2.7(16)}$	$5.7(14)^{+1.0(14)}_{-2.9(14)}$	<1.0(18)	<1.0(16)
G139.9091 4	$\frac{1.2(18)^{+1.0(17)}_{-4.7(17)}}{1.2(18)^{+5.2(17)}_{-4.7(17)}}$	$\frac{1.0(17)^{+2.0(10)}_{-4.3(16)}}{1.0(17)^{+2.0(10)}_{-4.3(16)}}$	$5.6(14)^{+1.9(14)}_{-3.9(14)}$	<1.9(14)	<1.0(16)
G075.78 1	$2.2(18)^{+3.2(17)}_{-2.9(17)}$	$3.1(17)^{+0.9(16)}_{-3.9(16)}$	$1.5(17)^{+2.4(10)}_{-1.7(16)}$	$2.0(16)^{+5.6(15)}_{-8.6(14)}$	$8.0(17)^{+5.4(10)}_{-6.2(16)}$
G075.78 2	$9.0(17)^{+2.0(17)}_{-7.3(16)}$	$5.6(17)^{+0.5(10)}_{-6.0(16)}$	$1.8(16)^{+4.6(15)}_{-3.1(15)}$	<1.4(14)	<1.1(16)
G075.78 3	$6.4(17)^{+2.9(17)}_{-7.4(16)}$	$2.4(17)^{+7.7(10)}_{-8.2(16)}$	$2.1(15)^{+0.6(14)}_{-6.6(14)}$	<3.2(14)	<1.1(17)
G0/5.78 4	$\frac{2.0(18)_{-5.7(17)}^{+0.1(17)}}{2.0(10)_{-5.7(17)}^{+4.8(17)}}$	$\frac{1.6(17)^{+0.1(10)}_{-3.5(16)}}{2.1(17)^{+9.1(16)}}$	$\frac{2.2(15)_{-5.6(14)}}{2.7(1.6)^{+3.9(15)}}$	<3.5(14)	<2.7(16)
IRAS 21078 1	$2.9(18)^{+1.0(17)}_{-1.8(17)}$	$3.1(17)^{+1.1(10)}_{-4.0(16)}$	$3.7(16)^{+2.0(15)}_{-2.0(15)}$	$6.2(16)^{+7.9(15)}_{-3.9(15)}$	$4.1(17)_{-1.3(17)}^{+1.2(17)}$
IRAS 21078 2	$1.7(18)^{+1.2(17)}_{-1.2(17)}$	$3.6(17)^{+0.1(16)}_{-6.1(16)}$	$1.3(16)^{+2.2(15)}_{-1.9(15)}$	$1.5(16)_{-3.7(15)}^{+1.8(15)}$	<1.3(16)
IRAS 21078 3	$1.8(18)^{+1.2(17)}_{-5.0(16)}$	$4.8(17)^{+1.0(10)}_{-5.3(16)}$	$9.5(15)^{+2.2(15)}_{-1.8(15)}$	$2.1(16)^{+1.0(15)}_{-3.0(15)}$	<1.5(16)
IRAS 21078 4	$9.6(17)^{+1.0(17)}_{-1.9(17)}$	$5.7(16)^{+7.6(16)}_{-3.5(16)}$	$6.7(14)^{+0.0(11)}_{-4.1(14)}$	<3.4(14)	<1.9(16)
IRAS 21078 5	$5.3(17)_{-1.4(17)}$	$6.5(16)_{-3.1(16)}$	$1.4(15)_{-3.6(14)}$	$1.7(15)_{-1.2(15)}$ $1.0(15)^{+3.0(15)}$	<1.2(16)
IRAS 21078 6	$9.2(17)_{-1.2(17)}^{+5.9(17)}$	$4.6(17)_{-6.5(16)}^{+1.4(17)}$	$2.8(15)_{-1.1(14)}^{+3.3(14)}$	$1.9(15)_{-1.4(15)}^{+1.4(15)}$	<1.1(16)
IKAS 210787	$8./(1/)_{-1.1(17)}$	$1.7(17)^{+1.1(17)}_{-6.4(16)}$	$1.9(15)_{-4.8(14)}^{+1.2(15)}$	<1.0(14) 1 0(15)+1.2(14)	<2.4(18)
IRAS 21078 8	$5.8(17)_{-1.5(17)}^{+1.3(17)}$	$1.7(17)_{-6.7(16)}^{+1.4(16)}$	$3.7(15)_{-9.0(14)}^{+8.6(14)}$	$1.0(15)_{-8.8(14)}^{-8.8(14)}$	<9.8(17)
IKAS 21078 9	$3.2(17)_{-5.3(16)}^{+1.5(17)}$	$8.5(16)^{+4.9(16)}_{-4.9(16)}$	$1.7(15)^{+0.0(14)}_{-4.8(14)}$	<1.9(14)	< 0.5(16)
IRAS 21078-10	$/.1(1/)_{-1.2(17)}^{+1.0(17)}$	$1.3(17)^{+0.1(10)}_{-3.6(16)}$	$/.6(15)^{+1.0(15)}_{-1.9(15)}$	$(1.9(14))_{-5.8(14)}$	<1.2(16)
IKAS 21078 11	$/.1(1/)_{-1.1(17)}^{+2.5(17)}$	$1.6(17)^{+2.2(10)}_{-5.2(16)}$	$4.0(15)^{+2.1(15)}_{-8.1(14)}$	<1.1(15)	<1.4(16)
IKAS 21078 12	$1.0(18)^{+3.0(17)}_{-4.8(17)}$	$1.5(17)_{-7.9(16)}^{+1.8(16)}$	$1.8(15)^{+0.2(14)}_{-6.7(14)}$	$2.3(15)_{-1.8(15)}$	<1.4(16)
IRAS 21078 13	$4.0(17)^{+5.1(17)}_{-7.0(16)}$	$3.0(16)^{+1.0(16)}_{-2.0(16)}$	$1.1(15)_{-2.3(14)}^{+0.5(14)}$	<2.2(14)	<1.0(19)
IRAS 21078-14	$1.7(18)_{-2.4(17)}$	$1.3(17)_{-5.6(16)}$	$1.2(15)^{+1.5(14)}_{-1.6(14)}$	<4.3(14)	<1.6(16)

Article number, page 44 of 81

Position	¹³ CO	C ¹⁸ O	SO	OCS	SO ₂
NGC7538 IRS9 1	$2.4(18)^{+6.3(17)}_{-3.3(17)}$	$2.2(17)^{+1.1(17)}_{-5.3(16)}$	$4.0(16)^{+4.3(15)}_{-4.7(15)}$	$2.1(16)^{+4.7(15)}_{-1.7(15)}$	$7.0(17)^{+1.7(17)}_{-2.0(17)}$
NGC7538 IRS9 2	$9.1(17)^{+2.2(17)}_{-2.1(17)}$	$1.2(17)^{+5.9(16)}_{-4.7(16)}$	$2.1(15)^{+6.8(14)}_{-5.2(14)}$	$1.9(15)^{+4.9(15)}_{-1.2(15)}$	<1.5(16)
NGC7538 IRS9 3	$6.6(17)^{+9.9(16)}_{-5.8(16)}$	$1.3(17)^{+9.8(16)}_{-4.0(16)}$	$1.8(15)^{+2.4(14)}_{-5.6(14)}$	<3.0(14)	<1.4(18)
NGC7538 IRS9 4	$1.6(18)^{+4.3(17)}_{-2.8(17)}$	$7.4(16)^{+4.1(16)}_{-1.6(16)}$	$2.7(14)^{+4.2(14)}_{-4.1(13)}$	<3.5(14)	<1.0(16)
NGC7538 IRS9 5	$6.8(18)^{+1.5(18)}_{-1.0(18)}$	$2.6(17)^{+6.0(16)}_{-6.7(16)}$	$1.1(15)^{+2.7(14)}_{-3.7(14)}$	<2.8(14)	<6.3(18)
NGC7538 IRS9 6	$8.0(17)^{+1.2(17)}_{-1.3(17)}$	$2.6(17)^{+1.8(17)}_{-6.3(16)}$	$2.1(15)^{+1.2(15)}_{-6.7(14)}$	$1.6(15)^{+3.4(15)}_{-9.7(14)}$	<2.2(16)
NGC7538 IRS9 7	$6.9(17)^{+3.3(17)}_{-1.3(17)}$	$2.2(17)^{+8.6(16)}_{-5.2(16)}$	$2.0(15)^{+5.1(14)}_{-4.1(14)}$	$1.6(15)^{+2.5(14)}_{-1.5(15)}$	<1.7(16)
NGC7538 IRS9 8	$4.6(17)^{+5.8(16)}_{-7.5(16)}$	$1.6(17)^{+2.5(16)}_{-5.6(16)}$	$8.7(14)^{+1.1(15)}_{-4.8(14)}$	<1.0(14)	<1.4(16)
S87 IRS1 1	$1.1(18)^{+2.2(17)}_{-1.0(17)}$	$1.7(17)^{+1.4(17)}_{-5.5(16)}$	$1.8(15)^{+1.7(15)}_{-8.7(14)}$	<1.1(15)	<3.8(16)
S87 IRS1 2	$5.4(17)^{+5.8(16)}_{-7.1(16)}$	$2.2(17)^{+1.3(17)}_{-1.1(16)}$	$5.8(14)^{+2.0(14)}_{-2.8(14)}$	<6.4(15)	<1.1(16)
S87 IRS1 3	$5.0(17)^{+9.3(16)}_{-4.4(16)}$	$1.7(17)^{+3.7(16)}_{-6.3(16)}$	$7.1(14)^{+6.3(14)}_{-2.4(14)}$	<2.5(14)	<1.3(16)
S87 IRS1 4	$6.8(17)^{+3.9(17)}_{-3.3(16)}$	$1.4(17)^{+9.1(16)}_{-3.9(16)}$	$8.4(14)^{+1.2(15)}_{-4.9(14)}$	<5.1(14)	<1.5(16)
S87 IRS1 5	$1.0(18)^{+5.3(16)}_{-1.1(17)}$	$1.1(17)^{+5.5(16)}_{-5.1(16)}$	$3.7(15)^{+1.1(15)}_{-1.6(14)}$	<3.8(14)	<2.7(16)
S87 IRS1 6	$1.9(18)^{+6.3(17)}_{-7.9(17)}$	$1.8(17)^{+3.8(16)}_{-5.7(16)}$	$6.6(14)^{+7.3(14)}_{-2.6(14)}$	<5.6(14)	<1.2(18)
S87 IRS1 7	$5.6(17)^{+9.9(16)}_{-5.7(16)}$	$1.4(17)^{+5.5(16)}_{-7.9(16)}$	$4.2(14)^{+3.1(14)}_{-8.9(13)}$	<1.1(14)	<2.4(16)
S87 IRS1 8	$3.9(18)^{+7.6(17)}_{-7.6(17)}$	$1.2(17)^{+1.8(17)}_{-5.6(16)}$	$6.3(14)^{+2.4(14)}_{-3.7(14)}$	<1.4(14)	<1.0(16)
S87 IRS1 9	$7.4(17)^{+9.6(16)}_{-1.2(17)}$	$1.6(17)^{+1.3(17)}_{-6.2(16)}$	$2.0(15)^{+9.0(14)}_{-8.1(13)}$	<1.3(14)	<1.6(18)
S106 1	$5.7(17)^{+1.5(17)}_{-1.2(17)}$	$3.8(16)^{+1.8(16)}_{-2.3(16)}$	<2.3(14)	<4.0(14)	<1.3(16)
S106 2	$1.3(18)^{+3.6(17)}_{-1.0(17)}$	$3.3(17)^{+1.6(17)}_{-5.1(16)}$	$1.6(16)^{+5.2(15)}_{-2.0(15)}$	$5.2(15)^{+2.4(15)}_{-1.6(15)}$	<2.7(16)
S106 3	$1.4(18)^{+2.2(17)}_{-1.3(17)}$	$7.6(17)^{+4.3(16)}_{-4.0(16)}$	$2.0(15)^{+1.4(15)}_{-6.9(14)}$	<1.6(15)	<5.7(16)
S106 4	$6.8(17)^{+1.3(17)}_{-1.3(17)}$	$2.5(17)^{+4.6(16)}_{-3.2(16)}$	$9.0(14)^{+4.8(14)}_{-7.4(14)}$	<3.8(14)	<2.5(16)
W3 H2O 1	$3.8(17)^{+1.3(17)}_{-8.1(16)}$	$5.3(16)^{+4.0(16)}_{-2.3(16)}$	$8.6(15)^{+6.0(15)}_{-2.6(15)}$	$3.2(16)^{+4.8(15)}_{-3.4(15)}$	$5.0(16)^{+1.7(16)}_{-1.5(16)}$
W3 H2O 2	$5.7(17)^{+5.3(16)}_{-4.9(16)}$	$1.7(16)^{+2.2(15)}_{-6.6(15)}$	$4.0(15)^{+9.8(14)}_{-9.4(14)}$	$1.3(16)^{+2.8(15)}_{-1.3(15)}$	$2.9(16)^{+2.1(16)}_{-1.2(16)}$
W3 H2O 3	$3.9(18)^{+1.1(17)}_{-2.5(17)}$	$4.4(17)^{+3.6(16)}_{-4.9(16)}$	$6.9(16)^{+1.1(16)}_{-5.4(15)}$	$4.7(17)^{+5.5(16)}_{-4.9(16)}$	$5.7(17)^{+1.6(17)}_{-7.1(16)}$
W3 H2O 4	$1.3(18)^{+1.3(17)}_{-1.2(17)}$	$3.9(17)^{+6.7(16)}_{-3.9(16)}$	$1.6(17)^{+1.6(16)}_{-2.1(16)}$	$8.5(17)^{+1.5(16)}_{-1.5(16)}$	$7.7(17)^{+6.8(16)}_{-7.6(16)}$
W3 H2O 5	$1.1(18)^{+3.0(17)}_{-2.4(17)}$	$8.1(17)^{+3.6(16)}_{-5.1(16)}$	$2.5(16)^{+3.3(15)}_{-2.9(15)}$	$1.4(17)^{+5.5(16)}_{-2.6(16)}$	<1.4(16)
W3 IRS4 1	$3.0(18)^{+1.6(17)}_{-2.8(17)}$	$1.4(17)^{+2.5(16)}_{-3.7(16)}$	$7.6(16)^{+5.5(15)}_{-2.2(15)}$	$2.0(16)^{+7.4(15)}_{-5.4(15)}$	$4.4(17)^{+2.8(16)}_{-1.8(16)}$
W3 IRS4 2	$7.5(17)^{+1.6(17)}_{-5.8(16)}$	$3.4(17)^{+3.4(16)}_{-5.5(16)}$	$1.7(15)^{+8.6(14)}_{-6.8(14)}$	<1.0(14)	<1.7(18)
W3 IRS4 3	$1.1(18)^{+2.1(17)}_{-7.9(16)}$	$5.0(17)^{+7.7(16)}_{-6.7(16)}$	$3.5(15)^{+2.6(15)}_{-3.0(14)}$	<9.0(14)	<1.5(16)
W3 IRS4 4	$1.5(18)^{+2.8(17)}_{-1.9(17)}$	$5.9(17)^{+4.0(16)}_{-1.1(17)}$	$1.3(15)^{+8.1(14)}_{-4.3(14)}$	<1.0(15)	<1.7(16)
W3 IRS4 5	$7.3(17)^{+3.2(17)}_{-2.9(16)}$	$5.5(17)^{+5.8(16)}_{-6.3(16)}$	$1.6(15)^{+8.8(14)}_{-2.7(14)}$	<1.6(15)	<2.1(16)
W3 IRS4 6	$1.2(18)^{+7.0(16)}_{-1.2(17)}$	$5.4(17)^{+3.8(16)}_{-8.3(16)}$	$1.2(15)^{+6.8(14)}_{-2.1(14)}$	<9.0(14)	<1.1(16)

Table E.1: continued.

Table E.2: Molecular column densities derived with XCLASS.

Position	DCN	H ₂ CO	HNCO	HC ₃ N	$HC_3N;v_7=1$
IRAS 23033 1	<1.5(13)	$7.3(15)^{+2.4(15)}_{-1.5(15)}$	<3.6(15)	<1.7(14)	<2.3(16)
IRAS 23033 2	$1.4(14)^{+7.7(13)}_{-2.9(13)}$	$2.9(16)^{+2.3(15)}_{-4.7(15)}$	$1.3(15)^{+7.3(14)}_{-6.3(14)}$	$9.9(14)^{+6.2(14)}_{-1.9(14)}$	<1.1(14)
IRAS 23033 3	$8.3(13)^{+1.4(13)}_{-8.0(12)}$	$6.6(16)^{+1.4(16)}_{-5.1(15)}$	$5.5(15)^{+2.7(15)}_{-1.5(15)}$	$3.3(15)^{+5.8(14)}_{-5.9(14)}$	$8.5(14)^{+9.5(14)}_{-4.5(14)}$
IRAS 23033 4	$7.2(13)^{+7.7(13)}_{-3.2(13)}$	$3.4(15)^{+1.2(15)}_{-9.2(14)}$	$3.6(16)^{+3.7(15)}_{-3.1(16)}$	$1.1(14)^{+2.3(14)}_{-6.4(13)}$	<8.4(14)
IRAS 23033 5	$1.3(14)^{+2.1(13)}_{-4.8(13)}$	$2.2(15)^{+1.1(15)}_{-1.3(15)}$	<8.3(14)	$5.4(13)^{+3.4(13)}_{-3.7(13)}$	<1.1(14)
IRAS 23151 1	$4.4(13)^{+9.9(13)}_{-3.4(13)}$	$3.8(16)^{+5.4(15)}_{-6.8(15)}$	$1.5(16)^{+2.7(15)}_{-1.9(15)}$	$4.3(14)^{+1.2(15)}_{-2.2(14)}$	<1.6(14)
IRAS 23151 2	$6.0(13)^{+9.2(13)}_{-4.1(13)}$	$9.1(15)^{+1.6(15)}_{-3.5(15)}$	$4.9(14)^{+7.5(14)}_{-3.0(14)}$	<1.0(13)	<3.9(15)
IRAS 23151 3	$2.1(13)^{+1.2(14)}_{-1.2(13)}$	$3.1(15)^{+9.7(14)}_{-6.2(14)}$	<2.2(14)	<3.0(13)	<1.1(15)
IRAS 23151 4	$2.8(13)^{+1.6(13)}_{-1.6(13)}$	$2.5(15)^{+5.9(14)}_{-9.8(14)}$	<1.7(14)	<3.1(13)	<6.4(15)
IRAS 23151 5	$7.1(13)^{+2.2(14)}_{-3.8(13)}$	$7.7(14)^{+1.4(15)}_{-2.7(14)}$	<1.1(14)	<3.2(13)	<3.3(15)

Notes. $a(b) = a \times 10^{b}$.

•

Position	DCN	H ₂ CO	HNCO	HC ₃ N	$HC_3N;v_7=1$
IRAS 23385 1	$9.8(13)^{+6.3(13)}_{-2.8(13)}$	$2.8(16)^{+4.7(15)}_{-2.8(15)}$	$4.1(14)^{+2.8(14)}_{-1.2(14)}$	$1.2(15)^{+2.7(14)}_{-3.1(14)}$	<1.6(14)
IRAS 23385 2	$2.3(13)^{+1.1(14)}_{-1.0(13)}$	$1.9(16)^{+6.0(15)}_{-4.6(15)}$	<7.8(14)	$9.1(13)^{+1.6(14)}_{-6.1(13)}$	<1.0(14)
IRAS 23385 3	$1.3(14)^{+8.6(13)}_{-3.7(13)}$	$2.1(16)^{+5.7(15)}_{-4.4(15)}$	<3.1(14)	$2.3(14)^{+1.5(14)}_{-8.5(13)}$	<9.7(15)
IRAS 23385 4	$4.3(13)^{+3.1(13)}_{-2.4(13)}$	$2.4(16)^{+7.2(15)}_{-4.7(15)}$	<7.9(14)	$9.0(13)^{+1.3(14)}_{-6.4(13)}$	<3.6(14)
IRAS 23385 5	$4.4(13)^{+1.0(14)}_{-2.1(13)}$	$1.2(16)^{+4.5(15)}_{-2.7(15)}$	<3.5(14)	$1.4(14)^{+9.6(14)}_{5.6(12)}$	<2.5(16)
IRAS 23385 6	$7.6(13)^{+2.1(13)}_{-6.5(12)}$	$1.3(16)^{+5.3(15)}_{-1.4(15)}$	<3.9(15)	$1.1(14)^{+2.5(14)}_{-8.6(12)}$	<9.7(15)
IRAS 23385 7	$3.1(13)^{+3.8(13)}_{-1.5(12)}$	$1.6(16)^{+1.1(16)}_{-5.7(15)}$	$7.4(15)^{+1.9(16)}_{-7.0(15)}$	$1.0(14)^{+1.7(14)}_{-8.0(12)}$	<1.0(14)
IRAS 23385 8	$5.7(13)^{+3.5(14)}_{-1.7(13)}$	$6.1(15)^{+2.1(15)}_{-1.9(15)}$	<6.9(14)	<4.1(13)	<1.5(16)
AFGL 2591 1	$\frac{-1.7(13)}{1.1(14)^{+4.6(13)}_{-3.2(13)}}$	$5.0(16)^{+7.9(15)}_{-3.8(15)}$	$1.8(16)^{+2.2(15)}_{-8.5(14)}$	$3.9(15)^{+4.2(14)}_{-3.8(14)}$	$9.9(15)^{+1.7(15)}_{8.2(14)}$
AFGL 2591 2	<1.0(13)	$1.4(15)^{+3.9(14)}_{-4.3(14)}$	<1.8(14)	<1.2(14)	$7.0(14)^{+8.1(14)}_{-5.3(14)}$
AFGL 2591 3	$4.7(13)^{+4.4(13)}_{-1.2(13)}$	$1.0(15)^{+1.0(15)}_{-2.1(14)}$	<1.0(14)	<1.5(13)	< 9.7(15)
AFGL 2591 4	$2.8(13)^{+1.1(14)}_{-1.8(13)}$	$2.4(15)^{+1.2(15)}_{-3.9(14)}$	<2.5(14)	$1.7(14)^{+6.1(14)}_{-8.3(13)}$	<1.0(14)
CepA HW2 1	$1.4(14)^{+4.1(13)}_{-6.4(13)}$	$5.4(17)^{+4.8(16)}_{-3.0(16)}$	$9.1(16)^{+1.9(15)}_{-1.8(15)}$	$7.6(15)^{+8.1(14)}_{-6.1(14)}$	$1.3(16)^{+6.4(14)}_{-6.1(14)}$
CepA HW2 2	$2.2(13)^{+1.1(13)}_{-1.9(13)}$	$8.6(16)^{+2.9(16)}_{-1.0(16)}$	$8.7(15)^{+9.9(14)}_{-2.4(15)}$	<2.8(14)	$6.4(14)^{+1.4(15)}_{-2.0(14)}$
CepA HW2 3	$9.2(14)^{+1.6(13)}_{-2.9(13)}$	$5.4(16)^{+1.2(16)}_{-6.8(15)}$	$5.5(15)^{+9.4(13)}_{-4.5(14)}$	$9.3(14)^{+4.4(14)}_{-2.1(14)}$	$9.9(14)^{+1.0(15)}_{-4.9(14)}$
CepA HW2 4	< 9.7(12)	<1.8(14)	<1.0(14)	<3.7(14)	<8.9(14)
CepA HW2 5	$3.7(13)^{+8.7(13)}_{-2.3(13)}$	$8.8(15)^{+2.8(15)}_{-7.8(14)}$	<1.0(14)	<3.4(13)	<9.7(15)
G084.9505 1	$1.1(14)^{+3.6(13)}_{-2.7(13)}$	$1.8(16)^{+6.5(15)}_{-1.3(15)}$	$8.9(14)^{+2.7(14)}_{-2.0(14)}$	$1.9(14)^{+7.9(13)}_{-1.5(14)}$	$1.0(14)^{+8.8(11)}_{-9.2(10)}$
G084.9505 2	$6.2(12)^{+3.2(12)}_{-5.1(12)}$	$7.6(14)^{+9.1(14)}_{-1.8(14)}$	<1.2(14)	<5.2(13)	<1.0(14)
G084.9505 3	$1.8(13)^{+4.6(13)}_{-1.4(13)}$	$3.4(15)^{+4.4(15)}_{-8.2(14)}$	<1.6(14)	<4.7(13)	<1.5(14)
G084.9505 4	$1.4(13)^{+6.9(13)}_{-5.0(12)}$	$1.3(15)^{+1.0(15)}_{-6.7(14)}$	$6.9(14)^{+7.1(14)}_{-5.1(14)}$	$4.1(14)^{+1.0(15)}_{-3.7(14)}$	<1.3(16)
G084.9505 5	$4.9(13)^{+6.6(13)}_{-3.5(13)}$	$6.7(14)^{+5.8(14)}_{-3.6(14)}$	<7.9(14)	<1.4(13)	<6.1(16)
G084.9505 6	$1.1(13)^{+1.3(12)}_{-9.6(12)}$	$2.3(15)^{+3.1(15)}_{-1.5(15)}$	<1.0(14)	<1.9(13)	<6.7(14)
G084.9505 7	$1.5(13)^{+6.9(12)}_{-1.3(13)}$	$6.0(14)^{+5.5(14)}_{-3.7(14)}$	<1.1(16)	<2.0(13)	<1.3(14)
G084.9505 8	$2.6(12)^{+3.8(12)}_{-1.2(12)}$	$7.7(14)^{+3.8(14)}_{-5.4(14)}$	<3.6(14)	<1.0(17)	<1.1(14)
G094.6028 1	$1.0(14)^{+6.0(13)}_{-1.5(13)}$	$4.1(16)^{+8.0(15)}_{-9.7(15)}$	$4.4(15)^{+3.2(14)}_{-6.7(14)}$	$2.3(15)^{+1.1(15)}_{-4.4(14)}$	$6.5(14)^{+8.0(14)}_{-5.0(14)}$
G094.6028 2	$6.9(13)^{+2.0(13)}_{-3.1(13)}$	$5.9(14)^{+5.4(14)}_{-8.9(13)}$	<1.6(16)	<2.5(13)	<1.0(14)
G094.6028 3	$3.6(13)^{+8.9(12)}_{-3.2(13)}$	$5.5(14)^{+9.8(14)}_{-1.6(14)}$	<2.8(14)	<1.0(13)	<5.7(14)
G094.6028 4	$5.2(13)^{+5.5(13)}_{-2.0(13)}$	$7.1(14)^{+6.1(14)}_{-2.8(14)}$	<1.5(16)	$7.6(13)^{+7.5(13)}_{-5.5(13)}$	<3.3(16)
G094.6028 5	$5.8(13)^{+1.8(14)}_{-2.6(13)}$	$5.5(14)^{+9.8(13)}_{-4.2(14)}$	$6.3(15)^{+5.8(16)}_{-5.4(15)}$	<2.2(13)	<1.2(15)
G094.6028 6	$2.8(13)^{+3.0(13)}_{-2.0(13)}$	$7.1(14)^{+6.5(14)}_{-2.9(14)}$	<1.0(14)	<1.0(13)	<1.2(14)
G094.6028 7	$1.5(13)^{+9.4(13)}_{-4.4(12)}$	$3.7(14)^{+2.0(14)}_{-2.4(14)}$	<3.6(14)	<2.6(13)	<2.4(14)
G094.6028 8	<1.9(12)	$1.2(14)^{+1.6(13)}_{-2.5(13)}$	<4.6(14)	<1.6(13)	<1.4(14)
G100.38 1	$2.7(13)^{+3.3(12)}_{-2.4(13)}$	$4.9(15)^{+1.8(15)}_{-1.3(15)}$	$9.1(14)^{+3.3(14)}_{-2.1(14)}$	<1.2(14)	<2.2(15)
G100.38 2	<6.5(12)	$4.3(14)^{+4.0(14)}_{-2.1(14)}$	<1.7(14)	<2.4(13)	<5.2(14)
G100.38 3	<1.0(13)	$3.1(14)^{+2.0(14)}_{-1.6(14)}$	<2.8(14)	<1.9(13)	<1.7(15)
G100.38 4	<9.0(12)	$1.2(15)^{+3.5(14)}_{-1.1(15)}$	<1.1(14)	<8.3(13)	<2.0(15)
G100.38 5	<7.1(12)	<1.1(14)	<2.0(14)	<4.9(13)	<2.9(14)
G108.75 1	$4.1(13)^{+3.7(14)}_{-2.2(13)}$	$6.6(15)^{+2.9(15)}_{-1.7(15)}$	<3.8(14)	$6.3(13)^{+1.5(14)}_{-1.5(13)}$	<3.5(14)
G108.75 2	$5.0(13)^{+2.3(13)}_{-4.4(13)}$	$3.5(15)^{+3.4(15)}_{-1.1(15)}_{+4.2(14)}$	<2.1(14)	<3.1(13)	<6.7(14)
G108.75 3	$2.0(13)^{+2.5(13)}_{-1.5(13)}$	$8.4(14)^{+4.5(14)}_{-4.5(14)}_{+3.1(15)}$	<1.0(17)	<1.3(13)	<1.6(14)
G108.75 4	$1.4(13)^{+5.7(15)}_{-9.0(12)}$	$1.7(15)^{+5.1(15)}_{-3.9(14)}$	<1.9(14)	<2.8(13)	<4.0(14)
G108.75 5	$5.2(13)^{+1.0(14)}_{-4.5(13)}_{-7.4(12)}$	$3.9(15)^{+2.4(14)}_{-2.6(15)}_{+1.0(15)}$	<3.1(14)	<1.3(13)	<1.9(14)
G108.75 6	$1.1(14)^{+7.4(13)}_{-6.8(13)}$	$8.5(14)^{+1.0(15)}_{-2.5(14)}$	<1.0(17)	$4.3(13)^{+1.2(14)}_{-1.5(13)}$	<1.5(14)
G138.2957 1	$3.6(13)^{+5.0(13)}_{-1.4(13)}$	$4.8(14)^{+5.2(14)}_{-2.9(14)}$	<1.2(15)	<7.4(13)	<1.0(14)
G138.2957 2	$7.8(13)^{+2.2(14)}_{-3.4(13)}$	$5.0(14)^{+5.7(14)}_{-2.5(14)}$	<5.1(14)	<3.1(13)	<1.5(14)
G138.2957 3	$1.1(14)^{+1.2(14)}_{-1.4(13)}$	$1.6(15)^{+1.5(15)}_{-4.0(14)}$	<2.9(14)	$6.6(13)^{+1.5(14)}_{-3.1(13)}$	<2.0(14)

Article number, page 46 of 81

Position	DCN	H ₂ CO	HNCO	HC ₃ N	$HC_{3}N;v_{7}=1$
G138.2957 4	$9.7(13)^{+9.6(13)}_{-8.5(13)}$	$4.3(14)^{+1.4(14)}_{-1.8(14)}$	$2.9(14)^{+6.1(14)}_{-5.2(13)}$	<1.9(13)	<5.4(16)
G138.2957 5	$1.8(13)^{+7.2(13)}_{-1.0(13)}$	$6.4(14)^{+3.4(14)}_{-1.9(14)}$	<2.2(15)	<2.1(13)	<1.4(14)
G138.2957 6	$4.8(13)^{+8.9(13)}_{-3.1(13)}$	$1.1(15)^{+1.8(15)}_{-5.2(14)}$	<4.3(14)	<1.4(13)	<2.4(16)
G138.2957 7	$1.2(14)^{+7.1(13)}_{-5.2(13)}$	$1.2(15)^{+1.5(15)}_{-4.6(14)}$	<1.0(17)	<5.4(13)	<1.0(14)
G138.2957 8	<1.3(13)	$9.4(14)^{+6.6(14)}_{-3.9(14)}$	<2.2(14)	<1.6(15)	<1.0(15)
G138.2957 9	$1.9(13)^{+5.4(13)}_{-1.3(13)}$	$1.5(15)^{+2.3(15)}_{-1.9(14)}$	<1.1(14)	<3.3(13)	<9.7(16)
G138.2957 10	<1.4(13)	$7.2(14)^{+7.9(14)}_{-8.8(13)}$	<1.1(14)	<1.7(13)	<3.8(14)
G138.2957 11	<7.2(12)	$1.7(15)^{+1.9(15)}_{-9.0(14)}$	<3.9(14)	<3.9(13)	<1.4(14)
G138.2957 12	$5.6(13)^{+3.5(13)}_{-4.8(13)}$	$7.1(14)^{+1.3(15)}_{-3.7(14)}$	<1.0(14)	<5.9(13)	<1.0(14)
G139.9091 1	<8.4(12)	$1.1(15)^{+1.3(14)}_{-9.6(14)}$	<1.1(14)	<7.4(13)	<6.5(14)
G139.9091 2	<2.2(13)	$1.3(15)^{+6.5(14)}_{-1.0(15)}$	<2.2(14)	<2.0(13)	<9.7(15)
G139.9091 3	$5.8(13)^{+2.5(14)}_{-3.0(13)}$	$3.6(14)^{+9.9(14)}_{-1.0(14)}$	<1.9(14)	<7.7(13)	<9.7(15)
G139.9091 4	<1.2(13)	$7.2(14)^{+5.8(14)}_{-5.1(14)}$	<2.9(14)	<1.0(13)	<1.8(14)
G075.78 1	$1.2(14)^{+2.6(13)}_{-3.1(13)}$	$5.8(16)^{+2.2(16)}_{-7.5(15)}$	$1.5(16)^{+1.1(15)}_{-1.8(15)}$	$6.2(15)^{+5.3(14)}_{-5.9(14)}$	$1.6(16)^{+7.7(14)}_{-5.5(15)}$
G075.78 2	$9.6(13)^{+7.3(13)}_{-2.1(13)}$	$1.9(15)^{+1.4(15)}_{-5.9(14)}$	<1.8(14)	<1.3(13)	<8.3(14)
G075.78 3	<6.3(12)	$7.3(14)^{+3.4(14)}_{-2.1(14)}$	<3.1(14)	<1.1(13)	<7.5(15)
G075.78 4	$1.3(14)^{+8.1(13)}_{-3.1(13)}$	$8.3(15)^{+2.2(15)}_{-1.0(15)}$	<1.1(14)	<1.2(13)	<3.6(14)
IRAS 21078 1	$9.4(13)^{+9.6(13)}_{-3.5(13)}$	$4.8(17)^{+3.2(16)}_{-3.0(16)}$	$4.1(16)^{+4.6(15)}_{-2.8(15)}$	$6.5(15)^{+1.7(15)}_{-1.1(15)}$	$4.3(15)^{+1.4(15)}_{-1.1(15)}$
IRAS 21078 2	$1.6(14)^{+1.4(14)}_{-2.0(13)}$	$4.4(16)^{+8.8(15)}_{-6.1(15)}$	$5.0(15)^{+2.5(15)}_{-1.0(15)}$	$3.7(14)^{+6.5(14)}_{-2.1(13)}$	$2.6(14)^{+3.0(14)}_{-8.7(13)}$
IRAS 21078 3	$9.8(13)^{+9.1(13)}_{-1.9(13)}$	$6.5(16)^{+1.2(16)}_{-1.2(16)}$	$1.2(15)^{+6.4(14)}_{-1.6(14)}$	$5.0(14)^{+2.4(14)}_{-1.6(14)}$	<4.7(14)
IRAS 21078 4	$4.1(13)^{+2.4(13)}_{-2.4(13)}$	$1.6(15)^{+1.3(15)}_{-2.5(14)}$	<1.0(14)	$1.2(14)^{+1.2(14)}_{-6.8(13)}$	<4.7(14)
IRAS 21078 5	$1.6(13)^{+1.6(13)}_{-1.3(13)}$	$2.1(15)^{+1.2(15)}_{-5.6(14)}$	<1.5(14)	<8.3(13)	<5.7(14)
IRAS 21078 6	$7.6(13)^{+4.2(13)}_{-3.2(13)}$	$1.2(16)^{+1.0(16)}_{-4.1(15)}$	<1.9(14)	$1.8(14)^{+2.2(14)}_{-8.9(13)}$	<9.7(15)
IRAS 21078 7	$1.3(14)^{+7.0(13)}_{-3.8(13)}$	$4.8(15)^{+1.2(15)}_{-1.4(15)}$	<5.2(14)	<9.1(13)	<9.4(14)
IRAS 21078 8	$1.3(14)^{+8.5(13)}_{-3.7(13)}$	$3.9(15)^{+1.8(15)}_{-7.2(14)}$	<2.5(14)	$1.8(14)^{+7.5(13)}_{-1.6(14)}$	<1.0(14)
IRAS 21078 9	$4.6(13)^{+6.1(13)}_{-2.0(12)}$	$5.2(15)^{+5.9(14)}_{-1.4(15)}$	<1.1(14)	$1.5(14)^{+5.1(13)}_{-1.0(14)}$	<4.5(14)
IRAS 21078 10	$5.0(13)^{+5.9(13)}_{-3.4(13)}$	$2.7(16)^{+2.0(16)}_{-5.7(15)}$	$7.9(14)^{+2.4(14)}_{-6.2(14)}$	$1.9(14)^{+3.0(14)}_{-8.9(13)}$	<8.7(14)
IRAS 21078 11	<8.5(14)	$4.0(15)^{+3.6(15)}_{-7.7(14)}$	<3.3(14)	<8.3(13)	<1.1(15)
IRAS 21078 12	$8.5(13)^{+6.3(13)}_{-2.6(13)}$	$1.1(16)^{+4.1(15)}_{-1.7(15)}$	<1.4(14)	$1.6(14)^{+1.1(14)}_{-1.4(14)}$	<1.0(14)
IRAS 21078 13	$1.3(13)^{+9.9(12)}_{-2.2(12)}$	$4.7(15)^{+1.3(15)}_{-4.7(14)}$	<8.4(14)	<1.1(14)	<1.3(14)
IRAS 21078 14	$2.1(13)^{+2.1(13)}_{-1.7(13)}$	$2.0(16)^{+6.9(15)}_{-4.3(15)}$	<1.9(14)	$9.4(13)^{+2.1(14)}_{-5.0(13)}$	<1.4(14)
NGC7538 IRS9 1	$1.1(14)^{+2.8(13)}_{-2.7(13)}$	$3.1(17)^{+5.5(16)}_{-3.2(16)}$	$9.8(16)^{+7.9(14)}_{-2.7(14)}$	$3.8(15)^{+4.1(15)}_{-1.0(15)}$	$4.7(15)^{+1.2(15)}_{-9.7(14)}$
NGC7538 IRS9 2	$4.6(13)^{+4.8(13)}_{-4.1(13)}$	$4.1(15)^{+1.5(15)}_{-1.1(15)}$	<1.9(14)	<2.0(13)	<3.6(14)
NGC7538 IRS9 3	$1.3(14)^{+5.6(13)}_{-3.9(13)}$	$4.3(15)^{+2.2(15)}_{-6.9(14)}$	<1.2(14)	$8.1(13)^{+5.1(13)}_{-7.0(13)}$	<1.8(14)
NGC7538 IRS9 4	$1.1(14)^{+5.3(13)}_{-2.5(13)}$	$1.2(15)^{+9.5(14)}_{-6.1(14)}$	<1.0(14)	<2.8(14)	<2.0(14)
NGC7538 IRS9 5	$5.9(13)^{+8.1(13)}_{-3.0(13)}$	$2.1(16)^{+7.6(15)}_{-5.6(15)}$	<1.6(14)	$1.0(14)^{+1.9(14)}_{-5.1(13)}$	<1.0(14)
NGC7538 IRS9 6	$5.7(13)^{+9.3(13)}_{-4.4(13)}$	$1.2(16)^{+1.5(15)}_{-1.6(15)}$	<6.6(14)	$2.5(14)^{+4.4(14)}_{-1.4(14)}$	<2.4(14)
NGC7538 IRS9 7	$5.0(13)^{+2.1(13)}_{-3.2(13)}$	$1.3(16)^{+5.4(15)}_{-3.8(15)}$	<7.8(16)	<9.1(13)	<1.1(16)
NGC7538 IRS9 8	$4.3(13)^{+4.7(13)}_{-2.0(13)}$	$1.4(15)^{+1.1(15)}_{-6.0(14)}$	<1.6(14)	$5.8(13)^{+1.1(14)}_{-3.3(13)}$	<2.2(14)
S87 IRS1 1	$2.7(13)^{+2.9(13)}_{-2.0(13)}$	$2.6(15)^{+3.3(15)}_{-1.3(15)}$	<3.2(16)	<1.5(13)	<2.4(14)
S87 IRS1 2	$7.7(13)^{+1.3(14)}_{-3.2(13)}$	$1.0(15)^{+1.3(15)}_{-1.9(14)}$	<3.6(14)	$6.7(13)_{-5.6(13)}^{+6.6(12)}$	<3.1(15)
S87 IRS1 3	<4.3(12)	$6.6(14)^{+6.7(14)}_{-2.5(14)}$	<1.0(14)	<2.0(13)	<2.9(14)
S87 IRS1 4	$2.0(13)^{+9.1(12)}_{-1.7(13)}$	$6.5(14)^{+1.9(14)}_{-2.0(14)}$	<2.7(14)	$1.2(14)^{+6.5(13)}_{-9.9(13)}$	<2.8(14)
S87 IRS1 5	<1.1(12)	$1.0(15)^{+1.6(15)}_{-2.1(14)}$	<3.3(16)	<1.7(13)	<3.1(14)
S87 IRS1 6	$1.1(14)^{+2.1(14)}_{-8.2(13)}$	$1.2(15)^{+1.7(15)}_{-5.3(14)}$	<1.0(14)	<2.2(13)	<1.4(14)
S87 IRS1 7	$9.7(13)^{+1.9(13)}_{-4.1(13)}$	$1.1(15)^{+1.1(15)}_{-4.1(14)}$	$1.5(14)^{+1.4(14)}_{-5.2(12)}$	$1.6(14)^{+2.0(14)}_{-7.9(13)}$	<1.4(14)

Table E.2: continued.

A&A proofs: manuscript no. paper_CORE_chemistry_arxiv

Position	DCN	H ₂ CO	HNCO	HC ₃ N	$HC_{3}N;v_{7}=1$
S87 IRS1 8	$8.5(13)^{+4.3(14)}_{-5.9(13)}$	$1.1(15)^{+1.2(15)}_{-3.0(14)}$	<3.1(14)	<1.8(13)	<2.3(16)
S87 IRS1 9	<1.2(13)	$2.5(15)^{+1.2(15)}_{-1.1(15)}$	<1.5(14)	<1.3(13)	<1.5(14)
S106 1	<2.2(12)	$3.0(14)^{+2.4(14)}_{-1.3(14)}$	<1.1(14)	<8.6(13)	<4.5(16)
S106 2	$1.9(14)^{+7.4(13)}_{-5.3(13)}$	$1.7(16)^{+3.8(15)}_{-2.6(15)}$	$2.2(15)^{+5.5(14)}_{-7.5(14)}$	$2.0(15)^{+1.0(15)}_{-5.7(14)}$	<6.5(14)
S106 3	$2.1(14)^{+9.0(13)}_{-2.4(13)}$	$2.8(15)^{+\overline{9.7(14)}}_{-7.1(14)}$	<5.4(14)	$2.3(14)^{+7.3(13)}_{-3.8(13)}$	<3.3(14)
S106 4	$1.3(14)^{+5.2(13)}_{-3.1(13)}$	$9.3(14)^{+4.4(14)}_{-5.6(14)}$	<1.0(14)	$8.2(13)^{+5.1(14)}_{-8.1(12)}$	<2.0(14)
W3 H2O 1	$1.2(14)^{+2.6(13)}_{-2.6(13)}$	$1.9(16)^{+5.2(15)}_{-3.4(15)}$	$2.4(15)^{+1.3(15)}_{-4.1(14)}$	$4.0(14)^{+6.0(14)}_{-1.1(14)}$	<7.1(14)
W3 H2O 2	$3.7(13)^{+2.2(13)}_{-1.6(13)}$	$1.0(16)^{+3.1(15)}_{-8.6(14)}$	$4.8(14)^{+2.5(14)}_{-3.4(14)}$	$2.2(14)^{+9.0(14)}_{-5.6(13)}$	$9.9(14)^{+5.3(15)}_{-1.8(14)}$
W3 H2O 3	$2.4(14)^{+2.8(13)}_{-1.7(13)}$	$1.3(17)^{+2.7(16)}_{-1.5(16)}$	$5.2(16)^{+2.1(14)}_{-3.9(15)}$	$8.8(15)^{+9.4(14)}_{-1.0(15)}$	$1.2(16)^{+1.2(15)}_{-1.1(15)}$
W3 H2O 4	$4.4(14)^{+3.2(13)}_{-2.0(13)}$	$2.2(17)^{+2.5(16)}_{-2.2(16)}$	$6.7(16)^{+1.6(15)}_{-2.9(15)}$	$6.9(15)^{+1.1(15)}_{-3.3(14)}$	$4.2(16)^{+1.0(16)}_{-1.2(16)}$
W3 H2O 5	$1.3(14)^{+8.9(13)}_{-1.6(13)}$	$5.7(16)^{+1.6(16)}_{-1.5(16)}$	$6.7(14)^{+1.8(15)}_{-4.4(14)}$	$3.1(15)^{+5.9(14)}_{-7.1(14)}$	<5.4(14)
W3 IRS4 1	$3.9(13)^{+8.1(13)}_{-3.2(13)}$	$4.0(16)^{+6.3(15)}_{-7.9(15)}$	$1.3(16)^{+1.8(15)}_{-8.9(14)}$	$1.9(15)^{+4.9(14)}_{-3.9(14)}$	$7.8(14)^{+1.7(15)}_{-3.6(14)}$
W3 IRS4 2	$8.4(13)^{+1.6(14)}_{-4.5(12)}$	$1.3(15)^{+4.5(14)}_{-4.7(14)}$	<8.7(16)	$2.5(14)^{+2.4(14)}_{-7.5(13)}$	<1.7(14)
W3 IRS4 3	$3.4(13)^{+1.8(14)}_{-1.2(13)}$	$1.2(15)^{+5.6(14)}_{-2.7(14)}$	<1.0(15)	$2.2(14)^{+3.0(14)}_{-8.0(13)}$	<1.7(14)
W3 IRS4 4	<5.9(12)	$2.1(15)^{+1.2(15)}_{-8.9(14)}$	<1.0(14)	<5.4(13)	<1.7(14)
W3 IRS4 5	$5.3(13)^{+4.0(13)}_{-3.0(13)}$	$2.4(15)^{+5.1(14)}_{-1.0(15)}$	<1.0(14)	$1.2(14)^{+1.5(14)}_{-7.3(13)}$	<6.7(14)
W3 IRS4 6	$3.4(13)^{+5.5(13)}_{-2.6(13)}$	$2.5(15)^{+2.2(15)}_{-1.0(15)}$	<6.1(14)	$7.8(13)^{+1.3(14)}_{-3.2(13)}$	<2.5(14)

Table E.2: continued.

Notes. $a(b) = a \times 10^{b}$.

Table E.3: Molecular column densities derived with XCLASS.

Position	$HC_{3}N;v_{7}=2$	CH ₃ OH	$CH_3OH; v_t=1$	CH ₃ CN
IRAS 23033 1	<1.0(17)	$1.5(16)^{+9.0(15)}_{-6.5(15)}$	<1.9(16)	<2.7(13)
IRAS 23033 2	<1.0(16)	$5.3(16)^{+5.3(15)}_{-1.1(16)}$	$7.5(16)^{+7.7(16)}_{-6.6(16)}$	$4.6(14)^{+6.9(13)}_{-3.3(13)}$
IRAS 23033 3	<5.5(16)	$8.5(16)^{+3.6(16)}_{-1.2(16)}$	$1.8(17)^{+6.0(16)}_{-6.4(16)}$	$1.8(15)^{+2.7(14)}_{-2.4(14)}$
IRAS 23033 4	<6.0(16)	$1.1(16)^{+2.9(16)}_{-7.0(15)}$	<1.3(16)	<5.0(13)
IRAS 23033 5	<2.3(16)	<8.6(14)	<5.9(15)	<3.1(13)
IRAS 23151 1	<1.0(16)	$1.8(17)^{+4.8(16)}_{-3.2(16)}$	$7.5(17)^{+3.7(17)}_{-2.5(17)}$	$1.2(16)^{+6.2(15)}_{-2.9(15)}$
IRAS 23151 2	<1.0(16)	$1.9(16)^{+2.5(16)}_{-1.2(16)}$	$1.6(17)^{+4.6(16)}_{-1.6(17)}$	$2.1(14)^{+\overline{1.8(14)}}_{-9.8(13)}$
IRAS 23151 3	<4.0(16)	$8.6(15)^{+2.6(16)}_{-4.5(15)}$	<5.4(15)	$9.8(13)^{+1.1(14)}_{-2.1(13)}$
IRAS 23151 4	<3.2(16)	$9.1(15)^{+1.8(16)}_{-8.8(15)}$	<5.1(16)	<1.2(14)
IRAS 23151 5	<6.1(16)	<1.3(15)	<1.5(18)	<7.2(13)
IRAS 23385 1	<3.8(16)	$2.6(16)^{+4.2(15)}_{-3.7(15)}$	$3.2(17)^{+2.5(18)}_{-2.5(17)}$	$4.1(14)^{+7.6(13)}_{-3.6(13)}$
IRAS 23385 2	<4.1(15)	$2.5(16)^{+9.3(15)}_{-6.2(15)}$	<2.9(16)	$7.2(13)^{+1.3(14)}_{-3.6(13)}$
IRAS 23385 3	<2.2(15)	$2.6(16)^{+2.3(16)}_{-1.0(16)}$	$3.2(16)^{+1.8(17)}_{-2.5(16)}$	$1.4(14)^{+2.0(14)}_{-4.4(13)}$
IRAS 23385 4	<1.4(15)	$2.6(16)^{+1.5(16)}_{-5.6(15)}$	$5.4(16)^{+9.0(16)}_{-5.1(16)}$	$1.0(14)^{+1.0(14)}_{-5.1(13)}$
IRAS 23385 5	<2.4(16)	$2.4(16)^{+2.1(16)}_{-8.3(15)}$	<5.1(16)	<9.4(13)
IRAS 23385 6	<1.5(16)	$2.1(16)^{+4.1(16)}_{-7.7(15)}$	$2.6(16)^{+4.1(15)}_{-2.5(16)}$	$9.8(13)^{+3.8(14)}_{-3.2(13)}$
IRAS 23385 7	<2.6(15)	$1.8(16)^{+1.4(16)}_{-2.4(15)}$	<1.9(16)	<1.1(14)
IRAS 23385 8	<2.1(15)	$1.6(16)^{+1.3(16)}_{-5.3(15)}$	<4.8(16)	<9.2(13)
AFGL 2591 1	$4.2(15)^{+7.2(14)}_{-5.8(14)}$	$2.0(17)^{+5.2(16)}_{-2.4(16)}$	$3.0(17)^{+6.8(16)}_{-6.7(16)}$	$1.6(16)^{+2.6(15)}_{-1.9(15)}$
AFGL 2591 2	<6.3(16)	<4.5(15)	<4.8(16)	<3.4(14)
AFGL 2591 3	<7.8(16)	<1.3(16)	<4.3(16)	<1.0(13)
AFGL 2591 4	<8.0(16)	$2.3(16)^{+3.0(16)}_{-2.2(16)}$	<3.0(16)	<1.6(13)
CepA HW2 1	$2.1(16)^{+3.5(15)}_{-5.2(15)}$	$3.6(17)^{+4.3(16)}_{-1.3(16)}$	$1.3(18)^{+4.6(17)}_{-9.4(16)}$	$1.4(16)^{+8.9(15)}_{-1.2(15)}$
CepA HW2 2	<6.6(15)	$2.2(16)^{+2.6(16)}_{-8.9(15)}$	<1.4(15)	$3.9(15)^{+1.3(15)}_{-7.0(14)}$
CepA HW2 3	<1.1(15)	$5.4(17)^{+8.6(16)}_{-3.3(16)}$	$5.6(18)^{+7.6(17)}_{-5.5(17)}$	$1.7(16)^{+4.0(15)}_{-2.2(15)}$
CepA HW2 4	<6.0(16)	<1.2(14)	<2.4(15)	<1.0(13)

Notes. $a(b) = a \times 10^{b}$.

Article number, page 48 of 81

Position	$HC_3N;v_7=2$	CH ₃ OH	$CH_3OH; v_t=1$	CH ₃ CN
CepA HW2 5	<1.0(16)	$6.8(16)^{+1.5(17)}_{-2.7(16)}$	$5.4(16)^{+3.9(16)}_{-5.2(16)}$	<1.2(13)
G084.9505 1	<1.0(16)	$4.6(16)^{+1.6(16)}_{-1.2(16)}$	$1.0(17)^{+1.5(17)}_{-7.4(16)}$	$5.8(14)^{+2.5(14)}_{-1.2(14)}$
G084.9505 2	<2.0(15)	$4.4(15)^{+2.8(16)}_{-3.6(15)}$	<1.3(15)	<1.1(13)
G084.9505 3	<1.1(16)	$5.3(15)^{+5.1(15)}_{-1.3(15)}$	<2.3(17)	<1.5(13)
G084.9505 4	<1.0(16)	$2.3(15)^{+1.7(15)}_{-1.5(15)}$	<1.2(17)	<1.7(13)
G084.9505 5	<1.0(16)	$2.1(15)^{+4.5(15)}_{-1.7(15)}$	<1.1(17)	<2.1(13)
G084.9505 6	<6.3(16)	$5.1(15)^{+5.6(15)}_{-2.7(15)}$	$1.4(16)^{+1.9(16)}_{-1.2(16)}$	<2.9(13)
G084.9505 7	<3.4(16)	<5.2(15)	<4.1(17)	<2.0(13)
G084.9505 8	<4.4(16)	<2.0(16)	<5.3(15)	<1.2(17)
G094.6028 1	<1.0(16)	$1.9(17)^{+5.0(16)}_{-1.4(16)}$	$1.1(18)^{+6.5(17)}_{-2.4(17)}$	$2.3(15)^{+3.2(14)}_{-4.0(14)}$
G094.6028 2	<5.5(15)	<7.8(15)	$2.8(16)^{+7.8(16)}_{-2.3(16)}$	<8.7(16)
G094.6028 3	<1.3(15)	<1.7(15)	<3.1(16)	<2.0(17)
G094.6028 4	<1.6(15)	<1.4(15)	<4.9(16)	<1.7(13)
G094.6028 5	<4.5(16)	<1.8(15)	<1.1(17)	<1.9(17)
G094.6028 6	<4.7(15)	<2.9(15)	<1.5(17)	<1.0(13)
G094.6028 7	<1.0(16)	<3.6(15)	$3.4(16)^{+9.6(16)}$	<5.3(13)
G094.6028 8	<1.5(15)	<1.4(15)	<2.3(16)	<2.3(13)
G100.38 1	<1.1(16)	$1.6(16)^{+4.1(16)}$	<4.7(15)	$2.1(14)^{+1.1(14)}_{-5.2(12)}$
G100 38 2	<2 4(16)	<10(15)	<1.1(16)	<73(13)
G100.38 3	<1.2(16)	<1.0(15)	< 4.5(15)	<13(13)
G100.38.4	<1.2(10)	< 2.8(15)	< 1.8(16)	< 5.4(16)
G100.38 4 G100.38 5	<1.0(15)	< 9.0(13)	$22(16)^{+2.8(16)}$	<3.4(13)
G108.75.1	<1.0(15)	$\frac{(9.0(11))}{1.2(16)^{+2.5(16)}}$	$\frac{2.2(10)-1.7(16)}{<3.8(17)}$	$78(13)^{+1.2(14)}$
C108.75.2	<1.0(15)	$1.2(10)_{-1.0(16)}$	< 5.0(17)	$(13)_{-5.3(13)}$
G108.75 2	<2.8(10)	< 8.0(17)	< 0.0(10)	<1.0(13)
G108.75 3	<1.5(16)	$2.2(15)^{+1.0(15)}_{-2.1(15)}$	<7.9(16)	<3.9(13)
G108.75 4	<1.2(15)	$5.7(15)^{+1.6(16)}_{-4.6(15)}$	<6.6(16)	<3.5(13)
G108.75 5	<1.5(15)	$3.7(15)^{+1.8(16)}_{-3.1(15)}$	<6.7(15)	<3.5(13)
G108.75 6	<1.2(15)	<1.5(16)	<2.1(16)	<8.4(16)
G138.2957 1	<1.6(15)	<1.0(19)	<1.2(16)	<1.3(13)
G138.2957 2	<1.0(17)	<3.3(15)	<4.5(16)	<1.9(13)
G138.2957 3	<8.5(16)	<6.1(15)	<2.6(16)	<1.9(13)
G138.2957 4	<1.5(15)	<2.3(15)	<6.0(16)	<1.1(14)
G138.2957 5	<6.1(16)	<2.5(15)	<2.3(16)	<8.5(17)
G138.2957 6	<1.3(16)	<3.8(15)	<8.6(15)	<1.5(15)
G138.2957 7	<2.1(15)	<1.3(16)	<1.3(17)	<2.3(13)
G138.2957 8	<6.8(16)	$1.3(16)^{+1.2(16)}_{-1.2(16)}$	<1.0(19)	<1.0(13)
G138.2957 9	<1.7(16)	<1.0(16)	<7.2(16)	<1.6(16)
G138.2957 10	<1.5(16)	<3.4(15)	<8.4(17)	<8.1(14)
G138.2957 11	<1.5(16)	<3.6(15)	<3.2(16)	<1.1(14)
G138.2957 12	<1.0(16)	<1.4(15)	<5.9(15)	<1.3(13)
G139.9091 1	<2.0(15)	<8.7(14)	<3.5(16)	<6.0(13)
G139.9091 2	<1.4(15)	<2.8(15)	<2.9(18)	<2.6(17)
G139.9091 3	<1.5(16)	<7.4(14)	<2.4(15)	<1.0(15)
G139.9091 4	<1.5(16)	<2.3(15)	<4.7(16)	< 6.2(13)
G075.78 1	$1.1(16)^{+3.4(15)}_{-1.8(15)}$	$1.8(17)^{+2.2(16)}_{-2.9(16)}$	$3.9(17)^{+2.4(17)}_{-5.7(16)}$	$8.2(15)^{+1.6(15)}_{-1.1(15)}$
G075.78 2	<1.0(16)	<2.1(15)	<2.3(15)	<1.0(13)
G075.78 3	<1.9(15)	<1.3(15)	<7.9(18)	<1.0(13)
G075.78 4	<4.8(15)	$1.3(16)^{+2.9(15)}_{-1.2(16)}$	<6.1(16)	<3.3(13)
IRAS 21078 1	<1.4(15)	$1.3(18)^{+1.4(17)}_{-7.0(16)}$	$2.3(18)^{+1.7(17)}_{-2.3(17)}$	$1.5(16)^{+2.2(15)}_{-1.6(15)}$
IRAS 21078 2	<1.0(16)	$1.9(17)^{+5.7(16)}_{-3.7(16)}$	$4.0(17)^{+5.1(16)}_{-4.5(16)}$	$2.5(15)^{+8.6(14)}_{-5.4(14)}$
IRAS 21078 3	$9.0(15)^{+4.2(16)}_{-3.1(15)}$	$6.6(17)^{+2.7(17)}_{-1.3(17)}$	$1.2(18)^{+1.2(17)}_{-2.9(17)}$	$4.0(15)^{+8.3(14)}_{-1.2(15)}$
IRAS 21078 4	<1.0(16)	<1.2(16)	<9.3(17)	<1.5(13)
IRAS 21078 5	<1.1(15)	$7.5(15)^{+4.0(15)}_{-1.5(15)}$	<6.0(16)	<9.0(13)

Table E.3: continued.

Notes. $a(b) = a \times 10^{b}$.

Article number, page 49 of 81

Position	$HC_3N;v_7=2$	CH ₃ OH	$CH_3OH; v_t=1$	CH ₃ CN
IRAS 21078 6	<1.2(16)	$3.1(16)^{+2.2(16)}_{-1.3(16)}$	<5.5(15)	$5.1(13)^{+4.4(13)}_{-1.7(13)}$
IRAS 21078 7	<1.0(16)	$7.1(15)^{+8.6(15)}_{-2.5(15)}$	<1.4(16)	<1.6(16)
IRAS 21078 8	<1.0(16)	$2.9(16)^{+6.6(15)}_{-1.3(16)}$	<8.8(15)	$1.5(14)^{+1.0(14)}_{-7.1(13)}$
IRAS 21078 9	<1.0(16)	$2.6(16)^{+2.5(16)}_{-9.4(15)}$	<6.1(18)	<2.6(13)
IRAS 21078 10	<7.2(16)	$2.5(16)^{+1.4(16)}_{-5.5(15)}$	$2.3(16)^{+8.3(15)}_{-2.1(16)}$	$2.1(14)^{+8.8(13)}_{-8.5(13)}$
IRAS 21078 11	<1.2(15)	$5.5(17)^{+8.1(17)}_{-4.8(17)}$	<4.4(16)	$1.5(14)^{+1.0(14)}_{-6.1(12)}$
IRAS 21078 12	<2.9(16)	$1.8(16)^{+1.2(16)}_{-4.2(15)}$	<3.5(16)	$1.5(14)^{+3.9(14)}_{-5.2(12)}$
IRAS 21078 13	<1.0(15)	$2.9(16)^{+3.4(16)}_{-1.6(16)}$	<3.7(16)	<2.9(14)
IRAS 21078 14	<5.4(15)	$2.6(16)^{+3.6(16)}_{-9.7(15)}$	<1.9(16)	<3.8(13)
NGC7538 IRS9 1	<5.2(15)	$9.1(16)^{+4.2(16)}_{-2.0(16)}$	$3.3(17)^{+2.4(17)}_{-1.2(17)}$	$8.7(15)^{+2.5(15)}_{-1.3(15)}$
NGC7538 IRS9 2	<7.3(16)	$5.2(15)^{+2.5(16)}_{-2.8(15)}$	<3.4(15)	<4.2(17)
NGC7538 IRS9 3	<1.3(16)	$7.6(15)^{+4.6(16)}$	<2.6(16)	<6.2(16)
NGC7538 IRS9 4	<3.0(16)	$3.4(15)^{+7.7(15)}$	<9.9(18)	<1.4(14)
NGC7538 IRS9 5	<6.6(16)	$9.1(15)^{+9.2(15)}$	< 9.9(15)	<5.1(13)
NGC7538 IRS9 6	<1.0(16)	$4.6(16)^{+3.9(16)}_{-4.7(15)}$	<2.1(16)	$1.3(14)^{+1.4(13)}$
NGC7538 IRS9 7	<3.2(16)	$3.5(15)^{+2.9(15)}_{-2.2(15)}$	<2.2(15)	<7.0(13)
NGC7538 IRS9 8	<4.0(16)	$4.4(15)^{+1.4(16)}_{-2.0(15)}$	<6.3(15)	<2.5(13)
S87 IRS1 1	<1.1(15)	<1.8(15)	<1.5(16)	<1.7(17)
S87 IRS1 2	<1.0(16)	<1.7(16)	<2.5(17)	<3.1(13)
S87 IRS1 3	<1.5(16)	<5.1(18)	<1.3(16)	<3.8(13)
S87 IRS1 4	<1.4(15)	$3.4(15)^{+8.4(15)}_{-2.9(15)}$	<5.1(16)	<2.8(13)
S87 IRS1 5	<1.5(16)	<1.4(14)	<2.2(18)	<9.0(13)
S87 IRS1 6	<1.3(15)	$2.7(15)^{+5.0(15)}_{-2.4(15)}$	<2.0(16)	<9.3(13)
S87 IRS1 7	<3.4(16)	<4.4(15)	<1.5(18)	<2.5(13)
S87 IRS1 8	<1.2(15)	<1.1(16)	$7.5(17)^{+5.5(17)}_{-7.4(17)}$	<1.0(13)
S87 IRS1 9	<2.9(16)	<1.4(16)	<1.8(16)	<1.0(18)
S106 1	<2.8(16)	<1.6(15)	<1.5(16)	<3.7(14)
S106 2	<3.8(15)	<3.4(16)	<2.2(16)	$4.1(14)^{+8.0(13)}_{-6.7(13)}$
S106 3	<2.1(16)	<1.4(16)	<1.1(17)	<2.6(14)
<u>\$106.4</u>	<1.0(16)	< 6.4(14)	$\frac{<8.3(15)}{1.1(12)+4.7(17)}$	<1.0(14)
W3 H2O 1	<1.8(16)	$2.0(18)^{+3.2(17)}_{-1.2(17)}$	$1.4(18)^{+4.7(17)}_{-1.7(17)}$	$2.2(15)^{+7.1(14)}_{-5.3(14)}$
W3 H2O 2	<1.9(15)	$1.1(17)^{+2.0(16)}_{-1.2(16)}$	$4.4(17)^{+2.5(17)}_{-2.1(17)}$	$5.4(14)^{+6.4(13)}_{-4.7(13)}$
W3 H2O 3	$1.2(16)^{+3.3(15)}_{-2.1(15)}$	$1.4(18)^{+8.5(16)}_{-4.4(16)}$	$3.1(18)^{+2.6(17)}_{-2.0(17)}$	$6.0(16)^{+1.0(16)}_{-1.0(16)}$
W3 H2O 4	$1.6(16)^{+3.0(15)}_{-1.1(15)}$	$2.9(18)^{+2.9(17)}_{-1.8(17)}$	$2.5(18)^{+7.5(16)}_{-1.3(17)}$	$1.5(17)^{+1.7(16)}_{-1.2(16)}$
W3 H2O 5	<4.5(16)	$4.1(16)^{+1.3(16)}_{-9.8(15)}$	<1.5(16)	$5.0(14)^{+7.9(13)}_{-5.7(13)}$
W3 IRS4 1	<3.3(16)	$1.8(17)^{+1.7(16)}_{-3.5(16)}$	$3.0(17)^{+1.4(17)}_{-4.6(16)}$	$6.7(15)^{+2.5(15)}_{-1.3(15)}$
W3 IRS4 2	<8.7(15)	$2.1(16)^{+1.1(17)}_{-1.9(16)}$	<2.0(18)	<2.2(13)
W3 IRS4 3	<2.5(15)	$1.7(16)^{+3.8(16)}_{-1.0(16)}$	<1.2(17)	<3.2(14)
W3 IRS4 4	<3.8(15)	<5.4(15)	<2.6(16)	<2.6(17)
W3IRS45	<3.1(16)	<1.6(15)	<2.1(18)	<1.3(14)
W3 IRS4 6	<2.3(15)	$5.0(15)^{+5.0(15)}_{-1.8(15)}$	<1.2(18)	$1.7(14)^{+1.2(14)}_{-8.3(13)}$

Table E.3: continued.

Appendix F: Observed spectra

Figure F.1 shows the observed spectrum and corresponding XCLASS fit for all 120 positions analyzed in Sect. 4.2.



Fig. F.1: *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 51 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 52 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 53 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 54 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 55 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 56 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 57 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 58 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 59 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 60 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 61 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 62 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 63 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 64 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 65 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 66 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 67 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 68 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 69 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 70 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 71 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 72 of 81


Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 73 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 74 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 75 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 76 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 77 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 78 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 79 of 81



Fig. F.1 (cont.): *Top panel:* Observed (black line) spectrum and XCLASS fit (red line) for all 120 analyzed positions. Fitted molecular transitions are indicated by green vertical lines. *Bottom panel:* Optical depth profile (blue line) of all fitted transitions for all 120 analyzed positions.

Article number, page 80 of 81

Appendix G: Model results

Table G.1 shows the best-fit chemical age τ_{chem} , χ^2 value, and percentage of well modeled molecules for each initial condition model (HMPO, HMC, and UCHII model) for each core.

Table G.1: MUSCLE results for all three initial condition models. The best-fit chemical age τ_{chem} , χ^2 value, and percentage of well modeled molecules Υ are shown for each initial condition model.

	HMF	PO mode	<u>.</u>	HMC model			UCHu model		
Core	τ_{chem}	v^2	Ϋ́	Tchem	v^2	Ϋ́	τ_{chem}	v^2	Ϋ́
	(yrs)	λ	(%)	(yrs)	λ	(%)	(yrs)	λ	(%)
IRAS 23033 1	33 574	0.339	64	73 271	0.346	64	97 675	0.336	64
IRAS 23033 2	18 0 19	0.525	64	49 046	0.589	64	104 065	0.589	55
IRAS 23033 3	18 705	0.517	64	50019	0.559	64	96415	0.567	64
IRAS 23151 1	19415	0.458	64	49 222	0.557	64	83 510	0.444	82
IRAS 23385 1	17 547	0.581	55	48 5 1 0	0.498	55	87 011	0.644	45
IRAS 23385 2	20729	0.545	55	48 5 1 0	0.602	45	95 268	0.465	55
AFGL 2591 1	17 884	0.508	64	54635	0.562	64	83 510	0.441	82
CepA HW2 1	18 705	0.602	55	48 5 30	0.605	55	83 510	0.478	64
CepA HW2 2	19415	0.459	73	48 812	0.616	45	88 141	0.454	73
G084.9505 1	16510	0.475	55	48 5 1 0	0.533	55	83 510	0.537	55
G094.6028 1	17 884	0.593	55	56611	0.521	64	83 784	0.536	55
G100.38 1	18 705	0.483	64	54 090	0.459	64	83 602	0.432	73
G108.75 1	20 0 1 1	0.485	73	48 602	0.608	55	87 011	0.610	55
G108.75 2	20729	0.319	73	60 2 68	0.325	73	106 070	0.296	73
G075.78 1	18 3 30	0.471	64	49 547	0.532	64	84 222	0.410	82
IRAS 21078 1	17 157	0.602	73	55 234	0.509	64	92 402	0.516	73
IRAS 21078 2	18 168	0.601	55	48 876	0.542	64	84 884	0.575	55
NGC7538 IRS9 1	17 884	0.502	73	71 070	0.521	73	84 292	0.447	82
S87 IRS1 1	16784	0.383	64	59 223	0.337	64	83 510	0.413	64
W3 H2O 3	17 547	0.590	55	48 876	0.579	45	83 510	0.532	64
W3 H2O 4	17 292	0.514	55	54 635	0.563	64	85 920	0.485	73
W3 IRS4 1	20 0 1 1	0.492	64	67 2 38	0.570	64	86415	0.469	82