

Organic molecules in the Galactic center[★]

Hot core chemistry without hot cores

M. A. Requena-Torres¹, J. Martín-Pintado¹, A. Rodríguez-Franco^{1,2}, S. Martín³,
N. J. Rodríguez-Fernández⁴, and P. de Vicente⁵

¹ Departamento de Astrofísica Molecular e Infrarroja, Instituto de Estructura de la Materia-CSIC C Serrano 121, 28006 Madrid, Spain

e-mail: requena@damir.iem.csic.es

² Escuela Universitaria de Óptica, Departamento de Matemática Aplicada (Biomatemática), Universidad Complutense de Madrid, Avenida Arcos de Jalón s/n, 28037 Madrid, Spain

³ Instituto de Radioastronomía Milimétrica Av. Divina Pastora 7, Local 20, 18012 Granada, Spain

e-mail: martin@iram.es

⁴ Observatoire de Bordeaux, L3AB (UMR 5804), CNRS/Université Bordeaux 1, BP 89, 2 rue de l'Observatoire, 33270 Floirac, France

e-mail: nemesio.rodriguez@obs.u-bordeaux1.fr

⁵ Observatorio Astronómico Nacional, Centro Astronómico de Yebes Apartado 148, 19080 Guadalajara, Spain

e-mail: p.devicente@oan.es

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ABSTRACT

Aims. We study the origin of large abundances of complex organic molecules in the Galactic center (GC).

Methods. We carried out a systematic study of the complex organic molecules CH₃OH, C₂H₅OH, (CH₃)₂O, HCOOCH₃, HCOOH, CH₃COOH, H₂CO, and CS toward 40 GC molecular clouds. Using the LTE approximation, we derived the physical properties of GC molecular clouds and the abundances of the complex molecules. The abundances of complex organic molecules in the GC are compared with those measured in hot cores and hot corinos, in which these complex molecules are also abundant.

Results. The CH₃OH abundance between clouds varies by nearly two orders of magnitude from 2.4×10^{-8} to 1.1×10^{-6} . The abundance of the other complex organic molecules relative to that of CH₃OH is basically independent of the CH₃OH abundance, with variations of only a factor 4–8. We find that both the abundance and the abundance ratios of the complex molecules relative to CH₃OH in hot cores are similar to those found in the GC clouds. However, hot corinos show different abundance ratios than observed in hot cores and in GC clouds. The rather constant abundance of all the complex molecules relative to CH₃OH suggests that all complex molecules are ejected from grain mantles by shocks. Frequent ($\sim 10^5$ years) shocks with velocities $> 6 \text{ km s}^{-1}$ are required to explain the high abundances in gas phase of complex organic molecules in the GC molecular clouds. The rather uniform abundance ratios in the GC clouds and in Galactic hot cores indicate a similar average composition of grain mantles in both kinds of regions. The Sickle and the Thermal Radio Arches, affected by UV radiation, show different relative abundances in the complex organic molecules due to the differentially photodissociation of these molecules.

Key words. astrochemistry – ISM: clouds – ISM: molecules – radio lines: ISM – Galaxy: center

1. Introduction

Gas-grain interaction in the interstellar medium (ISM) can have a large impact on its chemistry because of the desorption and/or depletion of molecules onto grains mantles. The gas phase abundance of molecules, which are believed to be efficiently formed on dust grains such as, methanol (CH₃OH), ethanol (C₂H₅OH), and formaldehyde (H₂CO), can be enhanced by orders of magnitude due to the ejection/evaporation of these molecules from grain mantles.

One of the interesting features of the gas-phase chemistry after ejection of these molecules is the fast conversion of these parent molecules into daughter molecules. The relative abundance of dimethyl ether ((CH₃)₂O) is expected to change by

three orders of magnitude in a short period of time after CH₃OH is ejected to gas phase (Millar et al. 1991; Caselli et al. 1993; Charnley et al. 1995; Horn et al. 2004). Basically all models of alcohol-driven chemistry have indicated that (CH₃)₂O and methyl formate (HCOOCH₃) will reach their largest abundances 10^4 – 10^5 years after the ejection of CH₃OH into gas phase. However, Horn et al. (2004) have shown that HCOOCH₃ formation in gas phase from CH₃OH is less efficient than previously considered in the models. This casts some doubts on the gas-phase production of HCOOCH₃, and they propose that this molecule is also formed on grain mantles. The gas phase/grain formation of (CH₃)₂O has been studied by Peeters et al. (2006), showing that the main path to form (CH₃)₂O is by gas-phase reactions and that its formation on grains is a minor source of the observed abundances. Other organic molecules like formic acid (HCOOH) can be produced in both gas phase and grain mantle chemistry (Liu et al. 2002).

[★] Table 7 is only available in electronic form at the CDS via anonymous ftp to cdsarc.u-strasbg.fr (130.79.128.5) or via <http://cdsweb.u-strasbg.fr/cgi-bin/qcat?J/A+A/455/971>

So far, large abundances of complex molecules have been detected in three different kinds of objects: hot cores associated with massive star formation (Ikeda et al. 2001), hot corinos associated with low-mass star formation (Bottinelli et al. 2006), and the Galactic center (GC) clouds (Martín-Pintado et al. 2001). Their physical properties are very different. While the hot cores and hot corinos are small ($\lesssim 0.1$ pc), hot (>100 K), and very dense ($\gtrsim 10^5$ cm $^{-3}$) condensations, the GC molecular clouds show averaged scales of 20–30 pc, kinetic temperatures ~ 50 –200 K, and an averaged density of $\sim 10^4$ cm $^{-3}$ (Güsten & Philipp 2004). The molecular gas in the inner region of our Galaxy presents a different chemistry than in the Galactic disk (Martín-Pintado et al. 1997). In the GC, widespread large abundances of grain-processed molecules like CH₃OH, C₂H₅OH, and silicon monoxide (SiO) are observed (Gottlieb et al. 1979; Minh et al. 1992; Martín-Pintado et al. 1997, 2001; Hüttemeister et al. 1998). It has been proposed that the sputtering of grains and grain mantles produced by widespread shocks with moderated velocities of ≤ 40 km s $^{-1}$ are the responsible of the “rich” chemistry observed in the GC clouds. The origin of the large scale shocks is so far unclear. These shocks could be produced by cloud-cloud collisions associated with the large-scale dynamics in the context of a barred potential (Hasegawa et al. 1994; Hüttemeister et al. 1998; Rodríguez-Fernández et al. 2006), by wind-blown bubbles driven by evolved massive stars (Martín-Pintado et al. 1999), or by hydrodynamic (HD) or magneto hydrodynamic turbulence (MHD) (Morris & Serabyn 1996). Since the formation of some complex organic molecules is believed to proceed in gas phase after the passage of the shocks and their abundances rapidly evolve, one could use the abundances of these molecules to gain insight into the chemistry of complex molecules in the ISM and to constrain the age of shocks in the GC (Martín-Pintado et al. 2001).

Following these ideas we made a systematic study of 40 molecular clouds in the GC region, between Sgr B2 and Sgr C, in complex molecules believed to be formed on grains like CH₃OH, C₂H₅OH, and HCOOCH₃, in gas phase from CH₃OH like (CH₃)₂O and in gas phase and/or grains like HCOOH and H₂CO. We also searched for the less abundant isomer of HCOOCH₃, the acetic acid (CH₃COOH). As tracers of the total column density of the molecular gas, we observed molecules like C¹⁸O, ¹³CO, and CS. Our systematic study shows that all the complex organic molecules present similar relative abundances with respect to CH₃OH, except for the regions where photodissociation could be important. Furthermore, the abundance and abundance ratios of these complex molecules in the GC are similar to those observed in hot cores, while hot corino abundances seem to have similar ratios for different objects. Our results suggest that all complex organic molecules have been ejected from grain mantles and that their abundances represent, in first approximation, the grain mantle composition.

2. Observations and results

The observations of all transitions of the complex organic molecules shown in Table 1 were carried out with the IRAM 30-m radio telescope at Pico Veleta (Spain). The data were obtained in different seasons between 1996 and 2003. The half-power beam width of the telescope was 24″, 17″, and 12″ for the 3, 2, and 1.3 mm bands. The receivers, equipped with SIS mixers, were tuned to single sideband with image rejections $\gtrsim 10$ dB. The typical system temperatures were 300, 500, and 900 K for the 3, 2, and 1.3 mm lines, respectively. We used two filterbanks of 256×1 MHz and one of 512×1 MHz as spectrometers. The

Table 1. Molecular line parameters.

Molecule	Transition	Frequency (MHz)	E_u/k (K)	$\mu_x^2 S$
CH ₃ OH.....	3 ₀ → 2 ₀ E	145 093.75	27.06	2.38
	3 ₋₁ → 2 ₋₁ E	145 097.47	19.52	2.11
	3 ₀ → 2 ₀ A+	145 103.23	13.94	2.38
	5 ₀ → 4 ₀ E	241 700.22	47.95	3.91
	5 ₋₁ → 4 ₋₁ E	241 767.22	40.41	3.75
¹³ CH ₃ OH ...	5 ₀ → 4 ₀ A+	241 791.43	34.83	4.94
	3 ₀₃ → 2 ₀₂ E	141 595.48	26.71	2.43
	3 ₋₁₃ → 2 ₋₁₂ E	141 597.06	19.21	2.16
C ₂ H ₅ OH	3 ₀₃ → 2 ₀₂ A+	141 602.53	13.60	2.43
	4 ₁₄ → 3 ₀₃	90 117.61	9.36	5.35
	7 ₀₇ → 6 ₁₆	104 487.26	23.26	8.61
HCOOCH ₃ ..	9 ₀₉ → 8 ₁₈	142 285.05	37.17	12.66
	7 ₂₅ → 6 ₂₄ E	90 145.69	19.69	17.00
	7 ₂₅ → 6 ₂₄ A	90 156.48	19.67	17.00
HCOOH	8 ₀₈ → 7 ₀₇ E	90 227.61	20.09	20.99
	8 ₀₈ → 7 ₀₇ A	90 229.63	20.07	20.99
	9 ₄₅ → 8 ₄₄ E	111 408.48	37.27	19.13
	9 ₁₈ → 8 ₁₇ E	111 674.10	28.15	23.12
	9 ₁₈ → 8 ₁₇ A	111 682.19	28.13	23.12
(CH ₃) ₂ O ^a	7 ₂₆ → 7 ₁₇	104 703.30	31.07	5.43
	7 ₀₇ → 6 ₁₆	111 783.01	25.26	6.80
CH ₃ COOH ..	4 ₂₂ → 3 ₂₁	90 164.25	23.53	5.78
	5 ₀₅ → 4 ₀₄	111 746.79	16.13	9.65
CS	8 _{*8} → 7 _{*7} A	90 246.25	20.30	43.20
	8 _{*8} → 7 _{*7} E	90 203.44	20.30	43.20
	10 _{*10} → 9 _{*9} A	111 548.53	30.50	54.80
	10 _{*10} → 9 _{*9} E	111 507.27	30.50	54.80
C ¹⁸ O	3 → 2	146 969.03	14.12	11.56
	5 → 4	244 936.64	35.28	19.84
C ¹³ O	1 → 0	109 782.17	5.28	0.01
	2 → 1	219 560.35	15.82	0.02
¹³ CO	1 → 0	110 201.35	5.30	0.01
H ₂ ¹³ CO	2 ₀₂ → 1 ₀₁	141 983.74	7.11	10.87

^a EE, AA, EA, and AE substates blended. Only the most intense transition is given.

velocity resolutions provided by the filter banks were 3, 2, and 1.3 km s $^{-1}$ for the 3, 2, and 1.3 mm bands, respectively. Spectra were calibrated using the standard dual load system. We used the antenna temperature scale (T_A^*) for the line intensities because the emission is rather extended and completely fills the beam.

Figure 1 shows the location of the sources we observed superimposed on the large-scale SiO map of the GC by Martín-Pintado et al. (1997). The sources were selected from the SiO maps of Martín-Pintado et al. (1997). Some of these sources had already been observed in C₂H₅OH by Martín-Pintado et al. (2001). We also included some clouds in Sgr C, the Thermal Radio Arches (TRA) (Serabyn & Güsten 1987), and the Sickle (Serabyn & Güsten 1991), where the C₂H₅OH emission was not detected by Martín-Pintado et al. (2001). The selected molecular clouds are spread over the region between Sgr C and Sgr B2. The positions of the sources are given in galactic and equatorial coordinates in Table 2. The nomenclature used to designate them is similar to that of Martín-Pintado et al. (2001): MC stands for Molecular Cloud followed by a G and the galactic coordinates, as recommended by the IAU. We grouped the sources in different regions as shown in Fig. 1 and Table 2.

Figures 2 and 3 show samples of line profiles for all observed molecules toward selected sources. Only for some sources like MC G–0.02–0.07 we did observe all the molecular lines. The line profiles of molecules such as CH₃OH are different from

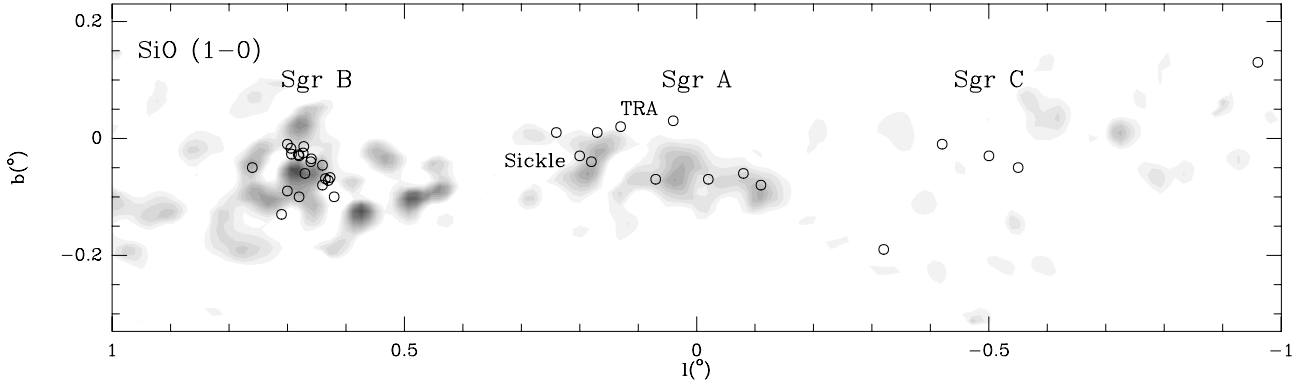


Fig. 1. Position of the observed sources superimposed on the SiO $J = 1 \rightarrow 0$ maps from the GC region by (Martín-Pintado et al. 1997).

Table 2. Source position.

Source	$l(^{\circ})$	$b(^{\circ})$	α (B1950)	δ (B1950)	Region
MC G-0.96+0.13	-0.96	+0.13	17 ^h 39 ^m 36 ^s .6	-29°39'47"	Sgr C
MC G-0.55-0.05	-0.55	-0.05	17 ^h 41 ^m 20 ^s .0	-29°24'30"	Sgr C
MC G-0.50-0.03	-0.50	-0.03	17 ^h 41 ^m 21 ^s .2	-29°21'26"	Sgr C
MC G-0.42-0.01	-0.42	-0.01	17 ^h 41 ^m 24 ^s .1	-29°15'51"	Sgr C
MC G-0.32-0.19	-0.32	-0.19	17 ^h 42 ^m 24 ^s .7	-29°17'20"	Sgr C
MC G-0.11-0.08	-0.11	-0.08	17 ^h 42 ^m 28 ^s .0	-29°02'55"	Sgr A
MC G-0.08-0.06	-0.08	-0.06	17 ^h 42 ^m 30 ^s .0	-29°00'58"	Sgr A
MC G-0.02-0.07	-0.02	-0.07	17 ^h 42 ^m 40 ^s .0	-28°58'00"	Sgr A
MC G+0.04+0.03	+0.04	+0.03	17 ^h 42 ^m 26 ^s .2	-28°51'45"	TRA
MC G+0.07-0.07	+0.07	-0.07	17 ^h 42 ^m 54 ^s .2	-28°53'30"	TRA
MC G+0.13+0.02	+0.13	+0.02	17 ^h 42 ^m 41 ^s .4	-28°47'35"	TRA
MC G+0.17+0.01	+0.17	+0.01	17 ^h 42 ^m 50 ^s .0	-28°45'50"	TRA
MC G+0.18-0.04	+0.18	-0.04	17 ^h 43 ^m 01 ^s .0	-28°47'15"	Sickle
MC G+0.20-0.03	+0.20	-0.03	17 ^h 43 ^m 03 ^s .6	-28°45'42"	Sickle
MC G+0.24+0.01	+0.24	+0.01	17 ^h 42 ^m 59 ^s .6	-28°42'35"	Sickle
MC G+0.62-0.10	+0.62	-0.10	17 ^h 44 ^m 18 ^s .0	-28°26'30"	Sgr B2_a
MC G+0.64-0.08	+0.64	-0.08	17 ^h 44 ^m 17 ^s .5	-28°24'30"	Sgr B2_a
MC G+0.67-0.06	+0.67	-0.06	17 ^h 44 ^m 18 ^s .0	-28°22'30"	Sgr B2_a
MC G+0.68-0.10	+0.68	-0.10	17 ^h 44 ^m 27 ^s .2	-28°23'20"	Sgr B2_a
MC G+0.70-0.01	+0.70	-0.01	17 ^h 44 ^m 10 ^s .0	-28°19'30"	Sgr B2_a
MC G+0.70-0.09	+0.70	-0.09	17 ^h 44 ^m 27 ^s .2	-28°22'05"	Sgr B2_a
MC G+0.71-0.13	+0.71	-0.13	17 ^h 44 ^m 38 ^s .4	-28°22'25"	Sgr B2_a
MC G+0.76-0.05	+0.76	-0.05	17 ^h 44 ^m 27 ^s .2	-28°17'35"	Sgr B2_a
SGR B2N	+0.68	-0.03	17 ^h 44 ^m 10 ^s .6	-28°21'17"	Hot Core
SGR B2M	+0.66	-0.04	17 ^h 44 ^m 10 ^s .6	-28°22'05"	Hot Core
MC G+0.694-0.017	+0.694	-0.017	17 ^h 44 ^m 10 ^s .0	-28°20'05"	Sgr B2
MC G+0.693-0.027	+0.693	-0.027	17 ^h 44 ^m 12 ^s .1	-28°20'25"	Sgr B2
MC G+0.627-0.067	+0.627	-0.067	17 ^h 44 ^m 12 ^s .1	-28°25'05"	Sgr B2
MC G+0.630-0.072	+0.630	-0.072	17 ^h 44 ^m 13 ^s .6	-28°25'05"	Sgr B2
MC G+0.672-0.014	+0.672	-0.014	17 ^h 44 ^m 06 ^s .1	-28°21'05"	Sgr B2
MC G+0.640-0.046	+0.640	-0.046	17 ^h 44 ^m 09 ^s .1	-28°23'45"	Sgr B2
MC G+0.635-0.069	+0.635	-0.069	17 ^h 44 ^m 13 ^s .6	-28°24'45"	Sgr B2
MC G+0.659-0.035	+0.659	-0.035	17 ^h 44 ^m 09 ^s .1	-28°22'25"	Sgr B2
MC G+0.681-0.028	+0.681	-0.028	17 ^h 44 ^m 10 ^s .6	-28°21'05"	Sgr B2
MC G+0.673-0.025	+0.673	-0.025	17 ^h 44 ^m 09 ^s .1	-28°21'25"	Sgr B2

Note. Galactic and equatorial coordinates of the selected sources. Nomenclature, as recommended by the IAU: MC followed by a G and the galactic coordinates, where MC stands for Galactic center molecular cloud. We have grouped the sources by regions: sources around Sgr C, Sgr A, the TRA, the Sickle, Sgr B2 (_a is to difference two observation sessions), and the hot cores. Not all sources in the TRA or in the Sickle are affected by the same physical conditions.

those of the CO isotopomers and CS due to the blending of several transitions. The complex organic molecules were only detected in some of the velocity components observed in C¹⁸O, ¹³CO, and CS. A good example of this difference in line profiles is shown in Figs. 2 and 3 for the source MC G+0.76-0.05

where C¹⁸O, ¹³CO, and CS show three or more velocity components, but C₂H₅OH only shows one. Each velocity component has been treated as an independent molecular cloud. Multiple velocity components are also found toward MC G+0.13+0.02, MC G+0.18-0.04, and MC G-0.32-0.19.

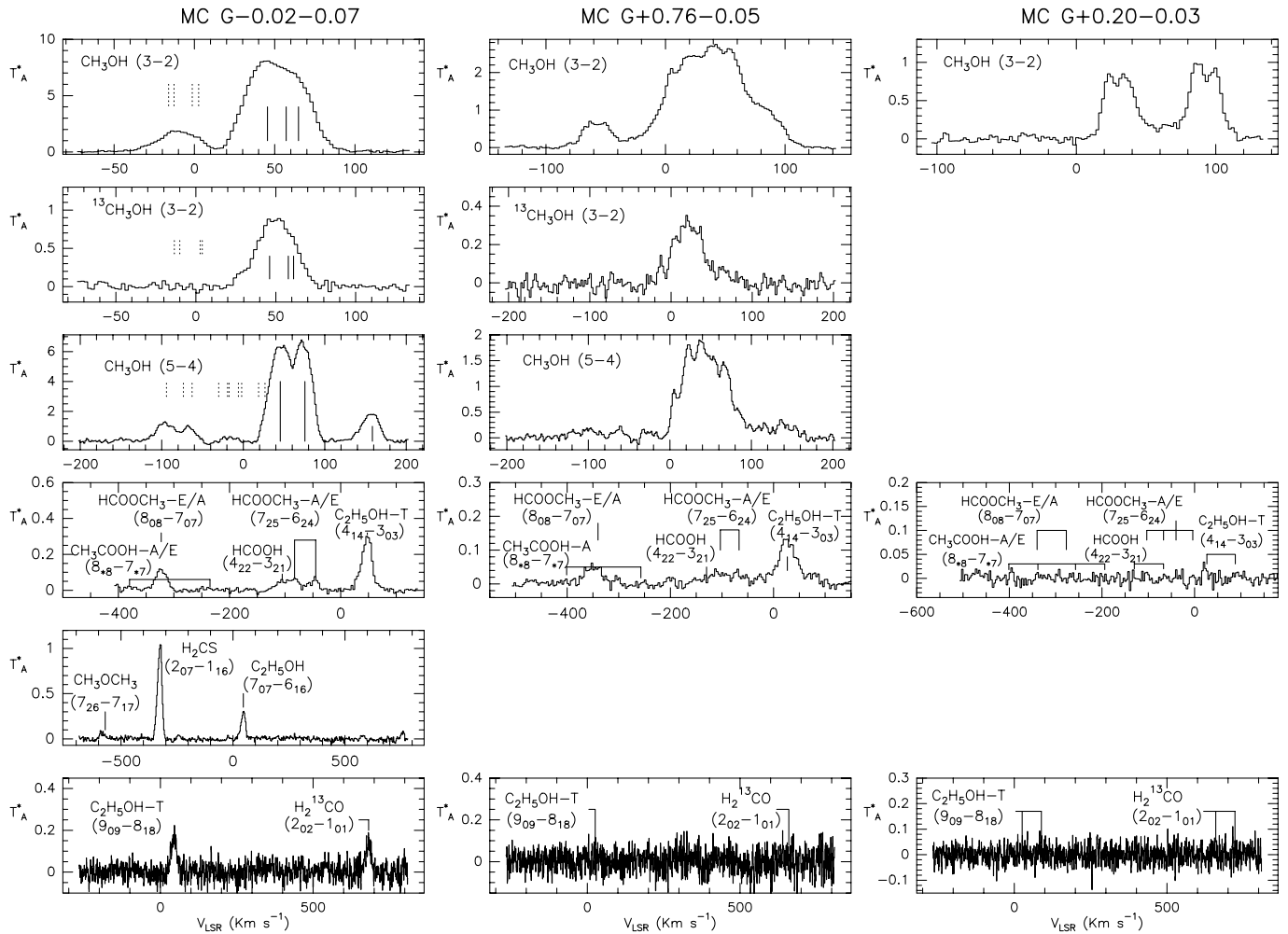


Fig. 2. Observed line profiles toward the sources MC G–0.02–0.07, MC G+0.76–0.05, and MC G+0.20–0.03. Line intensities are expressed in antenna temperature (K), and velocities refer to LSR (km s^{-1}). The CH_3OH profiles are composed of several overlapped transitions arising from levels with different energies; in the MC G–0.02–0.07 source, we show the transitions with highest intensities as vertical solid lines and the transitions with lowest intensities as dotted lines. These transitions are: for $\text{CH}_3\text{OH } J = 3 \rightarrow 2$, from left to right, $3_2 \rightarrow 2_2\text{A}$, $3_1 \rightarrow 2_1\text{E}$, $3_{-2} \rightarrow 2_{-2}\text{E}$, $3_2 \rightarrow 2_2\text{E}$, $3_2 \rightarrow 2_2\text{A}$, $3_0 \rightarrow 2_0\text{A}$, $3_{-1} \rightarrow 2_{-1}\text{E}$, and $3_0 \rightarrow 2_0\text{E}$. For the $^{13}\text{CH}_3\text{OH } J = 3 \rightarrow 2$, from left to right, $3_{21} \rightarrow 2_{20}\text{A}$, $3_{12} \rightarrow 2_{11}\text{E}$, $3_{21} \rightarrow 2_{21}\text{E}$, $3_{-22} \rightarrow 2_{-21}\text{E}$, $3_{22} \rightarrow 2_{21}\text{A}$, $3_{03} \rightarrow 2_{02}\text{A}$, $3_{-13} \rightarrow 2_{-12}\text{E}$, and $3_{03} \rightarrow 2_{02}\text{E}$. For the $\text{CH}_3\text{OH } J = 5 \rightarrow 4$ from left to right; $5_2 \rightarrow 4_2\text{E}$, $5_{-2} \rightarrow 4_{-2}\text{E}$, $5_2 \rightarrow 4_2\text{A}$, $5_1 \rightarrow 4_1\text{E}$, $5_{-3} \rightarrow 4_{-3}\text{E}$, $5_3 \rightarrow 4_3\text{E}$, $5_2 \rightarrow 4_2\text{A}$, $5_3 \rightarrow 4_3\text{A}$, $5_3 \rightarrow 4_3\text{A}$, $5_4 \rightarrow 4_4\text{E}$, $5_{-4} \rightarrow 4_{-4}\text{E}$, $5_4 \rightarrow 4_4\text{A}$, $5_4 \rightarrow 4_4\text{A}$, $5_0 \rightarrow 4_0\text{E}$, $5_{-1} \rightarrow 4_{-1}\text{E}$, and $5_0 \rightarrow 4_0\text{E}$. All these transitions have been used for the multi-Gaussian fit. The spectrum toward MC G+0.20–0.03 shows two velocity components.

3. Analysis of the data

To derive the physical and chemical properties of the selected molecular clouds, we fitted Gaussian profiles to all detected molecular lines for each velocity component. The observed spectra of some organic molecules are more complex than a single Gaussian profile because of the overlap of several transitions due to the large linewidths observed toward the GC clouds. To account for the overlap, we used the following constrains for the Gaussian fitting depending on the observed transitions:

- For molecules without internal rotation, like CS, C^{18}O , ^{13}CO , HCOOH, and H_2^{13}CO , and for molecules with internal rotation when the transitions are sufficiently separated, like those of $\text{C}_2\text{H}_5\text{OH}$ and some of the HCOOCH_3 lines, simple Gaussian profiles were fitted to each transition for every velocity component. As mentioned before, the CS, C^{18}O , and ^{13}CO profiles generally show more velocity components than those observed in the complex organic molecules. For this work we only fitted the components in the velocity range

where at least one organic molecule was detected. In the case where the velocity components overlapped we calculated the integrated intensity of CS, C^{18}O , and ^{13}CO in the velocity range where the complex organic molecules show the emission.

- The CH_3OH and $^{13}\text{CH}_3\text{OH}$ profiles are composed of several overlapped transitions arising from levels at different energies. In this case, we fitted the blended transitions with multi-Gaussian profiles forced to have the same linewidth and velocity separations corresponding to the rest frequencies of the transitions. We also forced the relative intensities derived from the spectroscopic parameters (Tables 1 and 3), assuming the same rotational temperature for all the transitions. In addition to the radial velocity and linewidth, we also fitted the rotational temperature and the optical depths that match the observed profiles. In this case, the antenna temperatures were obtained from $(T_{\text{rot}} - T_{\text{bg}})(1 - e^{-\tau})$. We used the optically thin emission from $^{13}\text{CH}_3\text{OH}$ transitions to test the optical depths and the antenna temperatures

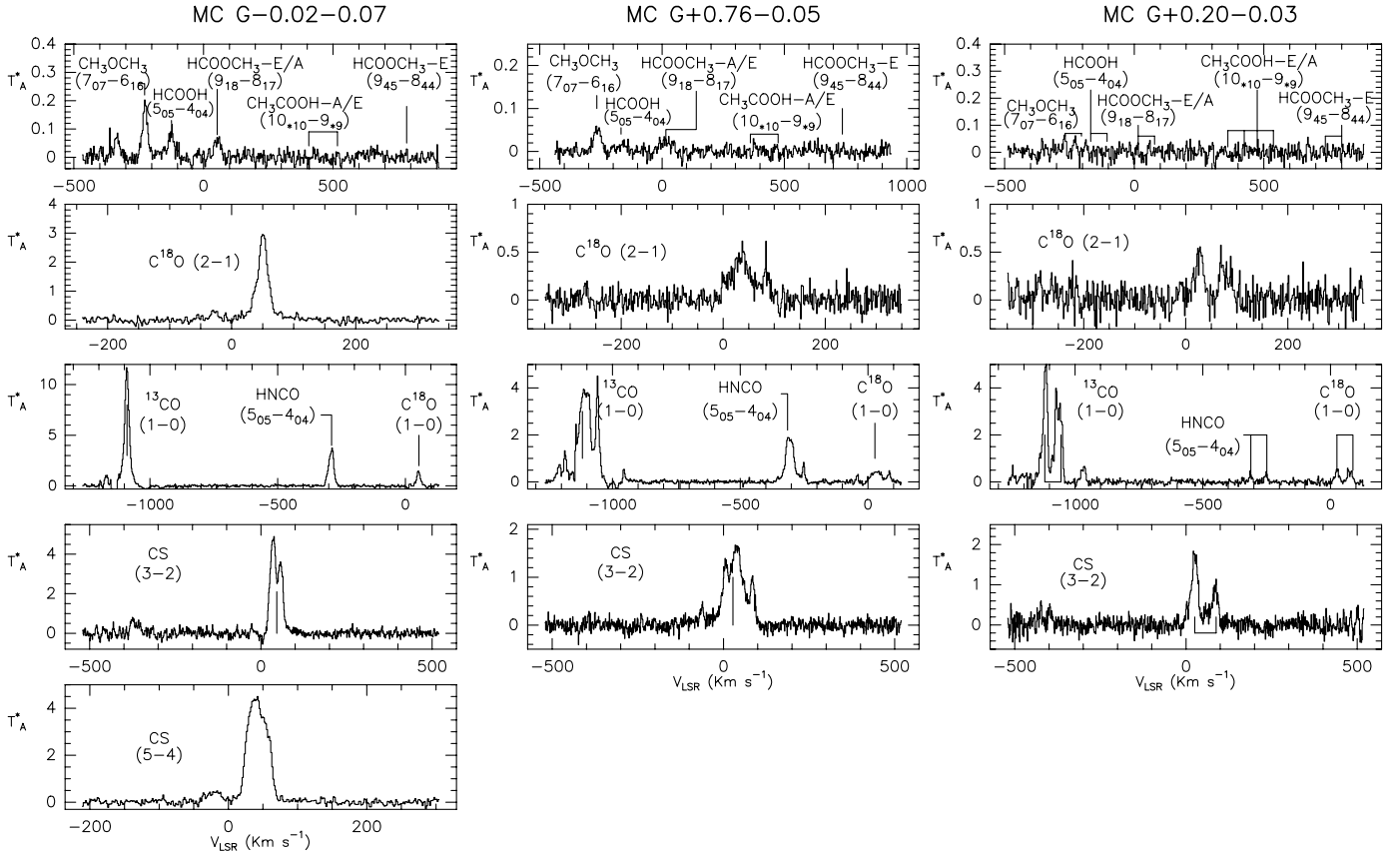


Fig. 3. Same as Fig. 2. The ^{13}CO , C^{18}O , and CS profiles show wider lines than those of the complex organic molecules in Fig. 2.

obtained from the CH_3OH fits. The optical depths derived from the CH_3OH lines are in good agreement with those obtained from the $[\text{CH}_3\text{OH}/^{13}\text{CH}_3\text{OH}]$ line-intensity ratios, assuming a $^{12}\text{C}/^{13}\text{C}$ ratio of 20 (Wilson & Rood 1994). For the $^{13}\text{CH}_3\text{OH}$, and in some cases for the CH_3OH lines, only the transitions with the largest intensities were fitted. In the case of the CH_3OH $5_0 \rightarrow 4_0\text{E}$ transition, a simple Gaussian fit was made since this line is not overlapped with any other transition.

- Due to its internal rotation, the $(\text{CH}_3)_2\text{O}$ rotational levels are split into four substates with similar Einstein coefficients but different intensities due to the nuclear spin degeneracy. These substates are called AA, EE, EA, and AE. Since the four substates are blended, we measured the integrated intensity of the four substates and then summed all the spin weights of each substate to derive the $(\text{CH}_3)_2\text{O}$ column densities.
- To obtain the upper limits to the column densities of undetected transitions like those of CH_3COOH , we used the 3σ level for the integrated intensities.

Derived parameters from the Gaussian fits (peak intensity, integrated intensity, radial velocity and linewidth) for all transitions in each source are shown electronically in Table 7, available at the CDS.

The linewidths of the complex organic molecules in our sources are $\sim 15 \text{ km s}^{-1}$, typical in the GC region. Radial velocities of the different sources range from -93 km s^{-1} to 140 km s^{-1} .

4. Derived parameters

4.1. Column densities, excitation temperatures, and densities

Molecular column densities were derived by assuming optically thin emission and the local thermodynamic equilibrium (LTE) approximation. In the case of optically thick lines like those of CH_3OH , the optically thin isotopic substitution was used. Under these conditions, the total column density, N , of a molecule is given by:

$$N = \frac{N_u}{g_u g_i} Q e^{(E_u/kT_{\text{rot}})} \quad (1)$$

$$N_u = 1.67 \times 10^{14} \frac{g_u}{\nu \mu^2 S} \int T_A^* dv, \quad (2)$$

where N_u is the column density in the upper level of the observed transition in cm^{-2} , g_u the upper level degeneracy ($2J_u + 1$), ν the line frequency in GHz, μ the dipole moment in Debye, S the line strength, $\int T_A^* dv$ the integrated intensity in K km s^{-1} , g_i the spin weight degeneracy, Q the partition function, E_u/k the upper level energy in K, and T_{rot} the rotational temperature also in K. Table 3 summarizes the dipole moments, the rotational constants, the partition functions, and the statistical weights used to calculate the total column densities.

4.1.1. Rotational temperatures

To derive the total column densities one first needs to estimate T_{rot} . For molecules with more than one observed transition, we calculated T_{rot} from population diagrams. Figure 4

Table 3. Dipole moments and rotational constants of the observed molecules.

Molecule	μ_a (Debye)	μ_b (Debye)	A (MHz)	B (MHz)	C (MHz)	Q	Comments	References
CH ₃ OH	0.885	1.440	127 484.00	24 679.98	23769.70	$1.28 T_{\text{rot}}^{1.5}$	Rotational levels split into E and A substates. Observed transitions with several blended lines.	Menten et al. (1986) Anderson et al. (1990a) Anderson et al. (1990b)
¹³ CH ₃ OH	0.899	1.440	127 527.40	24 083.50	23197.00	$1.28 T_{\text{rot}}^{1.5}$	Rotational levels split into E and A substates. Observed transitions with several blended lines.	Xu & Lovas (1997) Pearson et al. (1995, 1997)
C ₂ H ₅ OH	0.046	1.438	34 891.77	9350.68	8135.34	$3.2 T_{\text{rot}}^{1.5}$	Rotational levels split into Trans and Gauche substates. Only Trans Transitions are excited at ~10 K.	Plummer et al. (1984, 1986) Kuan & Snyder (1996) Groner et al. (1998)
HCOOCH ₃	1.630	0.680	19 985.71	6914.75	5304.48	$12.45 T_{\text{rot}}^{1.5}$	Rotational levels split into E and A substates.	Willemot et al. (1980) Liu et al. (2002)
(CH ₃) ₂ O	0.000	1.302	38 788.20	10 056.50	8886.80	$91.6 T_{\text{rot}}^{1.5}$	Rotational levels split into AA, EE, AE and EA substates. g_i : (K _a K _c : ee ↔ oo) AA:EE:AE:EA = 6:16:2:4 g_i : (K _a K _c : oe ↔ eo) AA:EE:AE:EA = 10:16:6:4	Włodarczyk & Demaison (1988) Mehring et al. (1997) Johnson et al. (1972)
HCOOH	1.396	0.260	77 512.25	12 055.11	10416.12	$1.68 T_{\text{rot}}^{1.5}$	Rotational levels split into E and A substates.	JPL JPL JPL
CH ₃ COOH	0.860	1.470	11 335.58	9478.73	5324.99	$14.1 T_{\text{rot}}^{1.5}$	Rotational levels split into E and A substates.	
H ¹³ CO	2.332		281993.04	37 809.11	33215.94	$0.57 T_{\text{rot}}^{1.5}$	Para (K _a = e) and Ortho (K _a = o) split, $g_i \rightarrow 1(p)-3(o)$.	
CS	1.957		24 495.56			$0.84 T_{\text{rot}}^{1.5}$		
C ¹⁸ O	0.111		54 891.42			$0.38 T_{\text{rot}}^{1.5}$		
¹³ CO	0.111		55 101.01			$0.38 T_{\text{rot}}^{1.5}$		

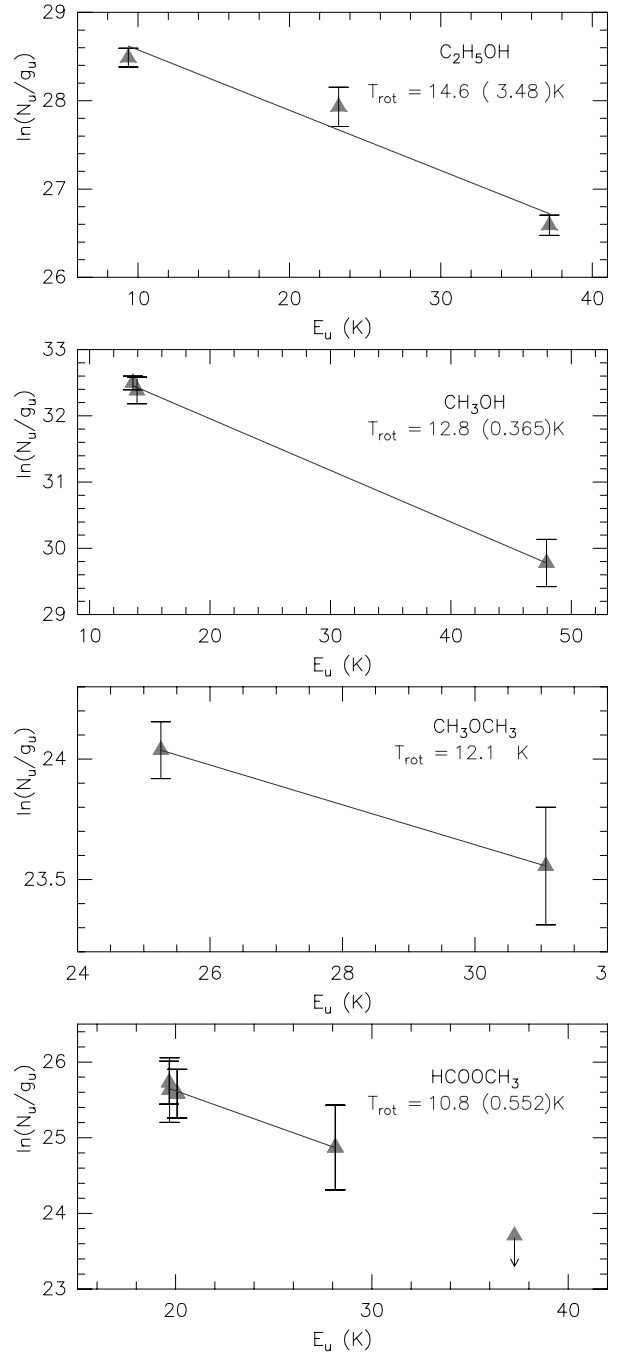


Fig. 4. Population diagrams for some complex molecules toward MC G–0.02–0.07. The error bars are derived by assuming a calibration uncertainty of 10% and the Gaussian fit error. Parentheses () indicate the calculated errors. Triangles with arrows represent upper limits to HCOOCH₃ column densities.

shows, as an example, the population diagrams ($\ln(N_u/g_u)$ vs. E_u/k) of some of the organic complex molecules derived for the source MC G–0.02–0.07. In general, T_{rot} from different molecules (third column on Table 4) agree within a factor of 2. We find T_{rot} values between 6 K and 18 K, much lower than the typical kinetic temperatures of ≥ 100 K derived from NH₃ by Hüttemeister et al. (1993) and H₂ by Rodríguez-Fernández et al. (2000, 2001a), indicating subthermal excitation. We do not find any systematic trend in T_{rot} as a function of the molecule. For sources with only one observed transition, we assumed an average value of $T_{\text{rot}} \sim 8$ K.

Table 4. Relative abundances and H₂ column densities.

source	V_{rad} (km s ⁻¹)	T_{rot} (K)	$X_{\text{CH}_3\text{OH}}$ $\times 10^{-7}$	$X_{\text{C}_2\text{H}_5\text{OH}}$ $\times 10^{-8}$	$X_{(\text{CH}_3)_2\text{O}}$ $\times 10^{-8}$	X_{HCOOCH_3} $\times 10^{-8}$	X_{HCOOH} $\times 10^{-9}$	$X_{\text{CH}_3\text{COOH}}$ $\times 10^{-9}$	$X_{\text{H}_2\text{CO}}$ $\times 10^{-9}$	X_{CS} $\times 10^{-9}$	$N(\text{H}_2)$ $\times 10^{22}$ cm ⁻²
MC G-0.96+0.13	140	8.0 ¹	10.6	≤1.4	≤8.3	≤10.7	≤12.2	≤30.0		16.6	0.2
MC G-0.55-0.05	-93	8.0 ¹	1.6	≤1.8	≤0.6	≤2.7	≤1.0	≤8.5		3.6	1.5
MC G-0.50-0.03	-92	8.0 ¹	0.4	≤1.0	≤0.3	≤1.1	≤0.4	≤3.6		1.6	2.5
MC G-0.42-0.01	-71	8.0 ¹	0.6	≤1.9	≤0.6	≤2.9	≤1.1	≤9.5		3.5	0.6
MC G-0.32-0.19	-70.4	8.0 ¹	0.9	≤0.8	≤18.1	≤6.2	≤23.4	≤17.5		10.0	0.2
MC G-0.32-0.19	-24	8.0 ¹	0.2	≤0.6	≤6.3	≤4.5	≤17.0	≤12.7		7.3	0.2
MC G-0.32-0.19	26	8.0 ¹	1.1	≤2.6	≤27.0	≤19.5	≤73.3	≤54.7		24.5	0.07
MC G-0.11-0.08	19.8	12.0 ² -10.0 ¹	11.3	6.0	5.6	7.5	3.7	≤3.9	19.5	25.0/43.4 ³	1.0
MC G-0.08-0.06	29.6	10.0 ² -12.0 ¹	4.0	2.5	4.7	≤2.6	4.1	≤4.5	7.1	19.0	0.6
MC G-0.02-0.07	47	10.8 ⁴ -14.0 ²	3.8	1.2	1.5	1.1	1.4	≤0.5	6.6	4.9/17.5 ³	6.8
MC G+0.04+0.03	-26	10.0 ¹	2.7	0.8	0.8	≤0.7	≤0.6	≤2.0	≤3.4	9.5	1.1
MC G+0.07-0.07	52	10.0 ¹ -12.8 ²	2.6	1.2	1.3	1.9	1.3	≤1.9	≤13.1	5.3	2.2
MC G+0.13+0.02	-30	8.0 ⁵	5.0	≤0.5	≤1.4	≤3.4	≤2.1	≤9.4		16.1	0.6
MC G+0.13+0.02	52	8.0 ⁵	2.4	≤0.5	≤1.6	≤3.7	≤2.4	≤10.5		9.6	0.4
MC G+0.17+0.01	59	10 ¹ -12.7 ²	1.5	0.4	≤0.3	0.5	1.2	≤0.9	≤3.1	3.7	1.3
MC G+0.18-0.04	27.1	8.0 ⁵	1.2	≤0.2	≤0.8	≤0.9	≤1.7	≤2.5	≤10.2	7.5/37.3 ³	0.7
MC G+0.18-0.04	85.8	8.0 ⁵	1.4	≤0.1	≤0.6	≤0.7	≤1.3	≤1.9	≤7.8	5.0/22.0 ³	0.8
MC G+0.20-0.03	24.9	8.0 ⁵	0.6	≤0.04	≤0.2	≤0.3	≤0.5	≤0.9	≤2.0	8.7	1.7
MC G+0.20-0.03	87.7	8.0 ⁵	1.5	≤0.09	≤0.4	≤0.7	≤1.1	≤2.0	≤4.4	8.3	0.9
MC G+0.24+0.01	36	12.6 ² -8.0 ¹	9.0	2.1	2.1	1.8	2.3	≤1.7	10.9	4.3/11.3 ³	3.0
MC G+0.62-0.10	56	11.0 ² -7.8 ¹	3.5	0.8	0.7	0.7	≤0.3	≤1.1	3.5	5.5	2.6
MC G+0.64-0.08	63	10.0 ²	10.0	0.7	≤0.3	0.7	≤0.4	≤1.8	≤2.4	3.8	2.0
MC G+0.67-0.06	51	10.5 ²	3.1	0.8	0.5	0.7	≤0.4	≤1.3	1.9	7.8	3.1
MC G+0.68-0.10	21	9.0 ² -6.0 ⁴	2.5	1.3	1.1	1.5	≤0.4	≤1.3	2.1	8.4	3.2
MC G+0.70-0.01	62	11.0 ² -10.0 ¹	3.7	0.9	1.5	1.1	1.2	≤1.6	4.2		2.8
MC G+0.70-0.09	43	10.0 ¹	1.7	0.7	≤0.3	≤1.3	≤0.4	≤3.6	≤2.4	5.3	2.9
MC G+0.71-0.13	40	8.0 ¹	2.4	≤0.8	≤0.6	≤2.8	≤0.9	≤7.9	≤0.6	6.0	2.8
MC G+0.76-0.05	27	10.0 ⁴	2.8	1.3	1.1	1.5	0.8	≤1.2	≤2.3	6.7	3.8
MC G+0.694-0.017	66	8.0 ¹ -11.0 ⁴	8.3	3.1	2.1	1.9	2.8	≤2.4		5.0	3.9
MC G+0.693-0.027	68	10.0 ¹ -11.5 ⁴	7.8	3.1	2.7	2.1	2.8	≤1.6		6.2/15.0 ³	4.1
MC G+0.627-0.067	49	10.0 ¹ -6.0 ⁴	4.8	1.4	2.2	≤0.8	1.6	≤2.2		6.0/11.7 ³	2.2
MC G+0.630-0.072	46	9.0 ¹ -12.0 ⁶	5.2	2.4	1.1	≤0.9	1.4	≤2.5		6.2	1.4
MC G+0.672-0.014	54	10.0 ¹	1.9	0.9	1.0	0.7	0.4	≤1.2		6.8	5.5
MC G+0.640-0.046	58	10.0 ¹ -7.0 ⁶	4.2	1.2	1.4	0.6	1.1	≤1.6		11.9	4.7
MC G+0.635-0.069	48	11.0 ⁶ -8.0 ⁴	4.6	0.9	1.1	1.6	≤0.6	≤2.5		6.7	2.3
MC G+0.659-0.035	62	11.0 ¹ -16.0 ⁴	4.2	0.8	1.1	0.9	1.3	≤0.7			13.9
MC G+0.681-0.028	65	11.0 ¹ -12.0 ⁴	2.9	1.0	1.9	2.5	3.9	≤0.8			8.8
MC G+0.673-0.025	68	18.5 ⁴ -13.9 ¹	2.4	0.6	0.9	1.2	1.8	≤0.8			14.6
SGR B2N	65	72.6 ²	1.3	2.3	3.5	5.0	9.9	≤0.7			19.0
SGR B2M	63	55.4 ²	0.3	0.3	0.3	0.4	0.4	≤0.2			65.0

¹ Derived from CH₃OH. ² Derived from C₂H₅OH. ³ From ¹³CS (Martín et al. 2006). ⁴ Derived from HCOOCH₃. ⁵ Assumed 8 K. ⁶ Derived from (CH₃)₂O.

4.1.2. Densities

Once the T_{rot} is derived, we can make a rough estimate of the H_2 densities, providing that the kinetic temperatures and the collisional cross section are known. The H_2 densities can be estimated from

$$n = \frac{A_{\text{ul}}}{\langle v\sigma \rangle_{\text{ul}}} \frac{1 + \left[1 - \exp\left(\frac{h\nu}{\kappa T_{\text{rot}}}\right) \right] \frac{1}{\exp\left(\frac{h\nu}{\kappa T_{\text{bg}}}\right) - 1}}{\exp\left(\frac{h\nu}{\kappa T_{\text{rot}}}\right) \exp\left(-\frac{h\nu}{\kappa T_{\text{k}}}\right) - 1} \quad (3)$$

where A_{ul} is the Einstein coefficient between the upper and the lower levels, $\langle v\sigma \rangle_{\text{ul}}$ the collisional rate coefficient between the same levels, ν the transition frequency, h the Plank constant, κ the Boltzmann constant, T_{rot} the rotational temperature, T_{bg} the background temperature, and T_{k} the kinetic temperature. Pottage et al. (2004) have calculated the collision rate coefficients ($\langle v\sigma \rangle$) between E- CH_3OH and para- H_2 for kinetic temperatures between 5 K and 200 K. Assuming a kinetic temperature of 100 K (Hüttemeister et al. 1993; Rodríguez-Fernández et al. 2001a), we obtain H_2 densities of $\sim 10^5 \text{ cm}^{-3}$ for the rotational temperature derived from CH_3OH of 6–18 K. Similar H_2 densities are also derived when the measured line intensities of E- CH_3OH are fitted using the large velocity gradient (LVG) approximation model, provided by J. Cernicharo, for the excitation of E- CH_3OH . We also estimated the H_2 densities from the excitation temperatures derived from $\text{C}_2\text{H}_5\text{OH}$. The collisional cross section in this case was estimated from those of CH_3OH but corrected for the difference in mass and size. These roughly estimated H_2 densities are also $\sim 10^5 \text{ cm}^{-3}$.

The derived H_2 densities are based on the collisional rates for a kinetic temperature of 100 K. However, the linewidths in the GC molecular clouds indicate the presence of supersonic turbulence and thus of non-equilibrium processes. In this case the collisional rates will not be described by the kinetic temperatures. However, for the observed transitions the rate coefficients do depend weakly on the increasing collisional velocities (Pottage et al. 2004), therefore our estimated densities will only be marginally affected by the effect of a larger velocity difference between the collisional partners than those represented by the kinetic temperature.

4.1.3. The H_2 column densities

To estimate the H_2 column density, we used the $J = 1 \rightarrow 0$ and $J = 2 \rightarrow 1$ C^{18}O and the $J = 1 \rightarrow 0$ ^{13}CO transitions using the LTE approximation. We used the isotopic ratios of $^{12}\text{C}/^{13}\text{C} = 20$, $^{16}\text{O}/^{18}\text{O} = 250$ (Wilson & Rood 1994), and a CO relative abundance of $\text{CO}/\text{H}_2 = 10^{-4}$ (Frerking et al. 1982). The last column in Table 4 shows the derived H_2 column densities. Our H_2 column densities, derived from the LTE approximation, are within a factor of ≤ 2 of those derived by Rodríguez-Fernández et al. (2001a) using the LVG approximation.

Table 4 also shows the relative CS abundance estimated from the CS column densities derived using the T_{rot} obtained from CH_3OH . Since our data refer to the line emission of the main CS isotopomer, optical depth effects might affect the derived CS column densities (Hüttemeister 1993). We compared our results with those derived from ^{13}CS by Martín et al. (2006) in several common sources. The CS column densities derived from ^{13}CS are always higher than those of the CS by a factor of 1–5. When the ^{13}CS determinations are available, Table 4 shows the two values estimated for the relative abundance of CS.

4.1.4. Column densities of complex organic molecules

To derive the column densities of the complex organic molecules we used the parameters in Table 3 and Eqs. (1) and (2). The relative abundance of complex molecules derived from their column densities and the H_2 column densities are shown in Table 4.

4.2. Abundances and abundance ratios

Table 4 shows the derived rotation temperatures, fractional abundances, and the H_2 column densities for the different molecular clouds. Different sources in the same line of sight are identified by their radial velocities. The fractional abundances of the complex organic molecules are very high, even higher than in the hot cores. The abundances we find reach values up to 1.1×10^{-6} for CH_3OH , 6×10^{-8} for $\text{C}_2\text{H}_5\text{OH}$, 5.6×10^{-8} for $(\text{CH}_3)_2\text{O}$, 7.5×10^{-8} for HCOOCH_3 , 4.1×10^{-9} for HCOOH , and 1.9×10^{-8} for H_2CO . We have only obtained upper limits to the abundance of CH_3COOH between $\leq 5 \times 10^{-10}$ and $\leq 5.5 \times 10^{-8}$.

There are some sources for which only upper limits to the abundances were measured for all the complex organic molecules except for CH_3OH . In some cases, like the sources with galactic longitude $\leq -0.30^\circ$, the upper limits to the abundances of all complex molecules are not relevant because of the lack of sensitivity of our observations. In this case the upper limits have not been included in the discussion. Other cases, like the $\text{C}_2\text{H}_5\text{OH}$ upper limits toward MC G+0.18–0.04, MC G+0.20–0.03 (Sickle), and MC G+0.13+0.02 (TRA) are relevant and therefore included in our discussion. The molecular emission toward the Sickle shows two velocity components at $\sim 25 \text{ km s}^{-1}$ and $\sim 80 \text{ km s}^{-1}$, which are believed to be affected by the UV radiation produced by the Quintuplet cluster (Rodríguez-Fernández et al. 2001b). The molecular emission toward MC G+0.13+0.02, located in the TRA, also seems to be affected by the UV radiation from the Arches Cluster. In fact only the -30 km s^{-1} velocity component, which is clearly associated with the ionized gas (Serabyn & Güsten 1987), shows lower $\text{C}_2\text{H}_5\text{OH}$ abundance than in other sources.

Because CH_3OH is the only molecule that has been detected in all sources and it is believed to play a central role in the formation of some of the observed complex organic molecules, we compared the abundances of the rest of the complex molecules with that of CH_3OH . Figure 5 summarizes the results illustrating the abundance of all the observed molecules relative to that of CH_3OH as a function of the CH_3OH abundance. As already mentioned, we have only included the significant upper limits in this figure. The main results are:

- In spite of the CH_3OH 's abundance changing by a factor of ~ 50 , all the complex organic molecules, except that of the CH_3COOH with only upper limits to its abundance, show a surprisingly constant relative abundance with respect to CH_3OH .
- The mean $[\text{C}_2\text{H}_5\text{OH}/\text{CH}_3\text{OH}]$ abundance ratio of $\sim 3.6 \times 10^{-2}$ is constant within a factor 3.7. The Sickle and the TRA clouds show upper limits to the $[\text{C}_2\text{H}_5\text{OH}/\text{CH}_3\text{OH}]$ ratios, which are a factor of ≥ 6 times lower than the mean ratio found in the typical GC clouds, a factor of ≥ 1.6 higher than the scattering of the data.
- The abundance ratios $[(\text{CH}_3)_2\text{O}/\text{CH}_3\text{OH}]$ and $[\text{HCOOCH}_3/\text{CH}_3\text{OH}]$ present similar mean values of $\sim 3.9 \times 10^{-2}$. These ratios show a somewhat larger dispersion than that of $\text{C}_2\text{H}_5\text{OH}$ but are constant within a factor of 7.
- For the $[\text{HCOOH}/\text{CH}_3\text{OH}]$ ratio, we obtained a mean value of $\sim 7.9 \times 10^{-3}$ and a dispersion of a factor of 5.3.

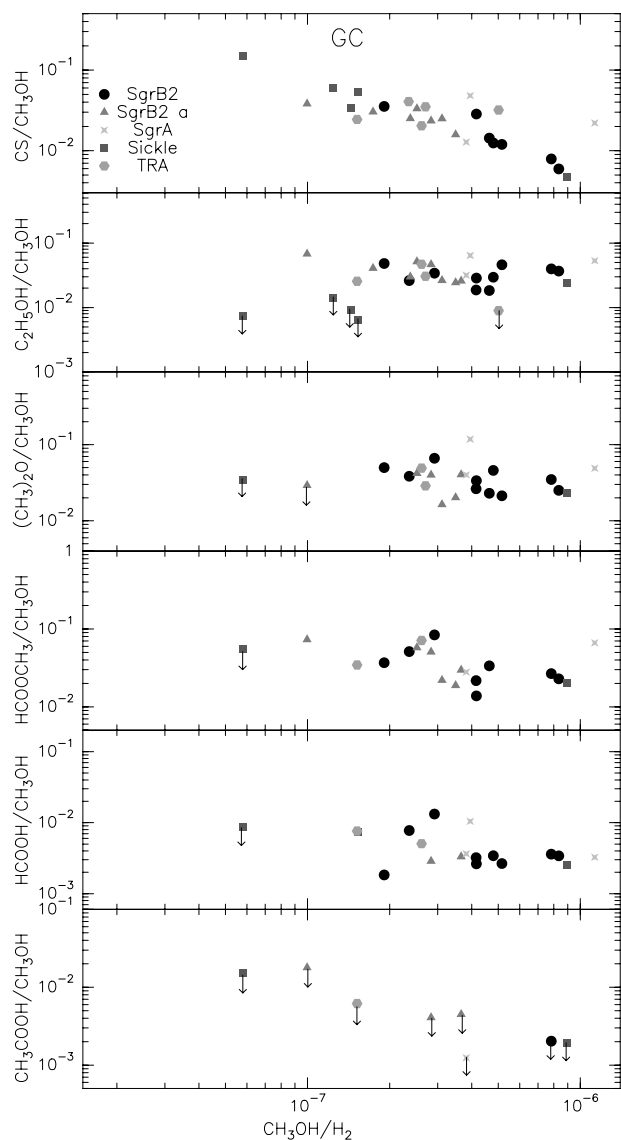


Fig. 5. Relative abundances of organic complex molecules with respect to CH_3OH as a function of the CH_3OH abundance for the selected GC clouds. Different regions are represented with different symbols (see Table 2). The arrows represent upper limits. Only the detections and significant upper limits are shown.

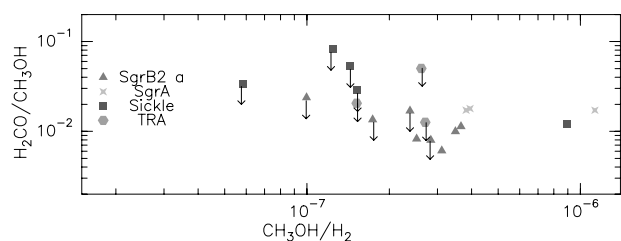


Fig. 6. Same as Fig. 5 but for H_2CO .

- CH_3COOH was not been detected in the GC clouds but its $[\text{CH}_3\text{COOH}/\text{CH}_3\text{OH}]$ ratio must be lower than 10^{-3} .
- We also show in Fig. 6 the $[\text{H}_2\text{CO}/\text{CH}_3\text{OH}]$ ratio as a function of the CH_3OH abundance. As for the more complex organic molecules, we find a constant ratio of $\sim 1.1 \times 10^{-2}$, within a factor 3.

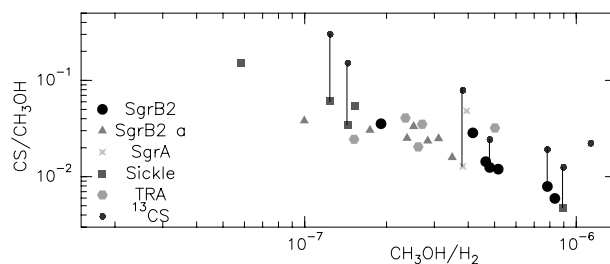


Fig. 7. Variation of $[\text{CS}/\text{CH}_3\text{OH}]$ as a function of CH_3OH abundances. The vertical bars show the changes due to the opacity.

The $[\text{CS}/\text{CH}_3\text{OH}]$ abundance ratio is shown in Fig. 7. This abundance ratio decreases by a factor 60 as the CH_3OH abundance increases. This is the only molecule for which we find a systematic trend, and the variation is clearly larger than those found for the complex organic molecules. As previously mentioned, our CS column densities could be affected by opacity effects. To show the effects of opacity, we have also included the abundance ratio obtained using the rarer CS isotopomer ^{13}CS in Fig. 7 as vertical bars. The $[\text{CS}/\text{CH}_3\text{OH}]$ abundance ratio derived from the optically thin isotopomer increases by a relatively constant value for all clouds independently of the CH_3OH abundance. The variation observed in the $[\text{CS}/\text{CH}_3\text{OH}]$ ratio as a function of the CH_3OH abundance, using the ^{13}CS data, is similar or even larger than that derived from the CS data. We therefore consider that the difference between the behavior of CS and the complex organic molecules is real and related to the different type of chemistry in the GC clouds.

5. Comparison with Galactic disk objects

In dark clouds, where the kinetic temperature is only 10 K, the abundance of complex organic molecules is very low. In the dark cloud TMC-1, the CH_3OH , H_2CO , and HCOOH abundances are as low as 2×10^{-9} , 2×10^{-8} , and 3×10^{-10} , respectively (see e.g. Turner 2000). For other dark clouds like L183, one finds similar abundances of 8×10^{-9} for CH_3OH , of 2×10^{-8} for H_2CO , and of 3×10^{-10} for HCOOH (Turner et al. 1999; Dickens et al. 2000). Other complex molecules, like $\text{C}_2\text{H}_5\text{OH}$, $(\text{CH}_3)_2\text{O}$, CH_3COOH , and HCOOCH_3 , have not been detected so far, but model predictions suggest $\text{C}_2\text{H}_5\text{OH}$ abundances of 10^{-12} (Hasegawa & Herbst 1993). High abundance of the complex organic molecules were first observed in the short-lived (10^{5-6} years) objects associated with massive star-forming regions known as hot cores. There, it is believed that grain mantle evaporation release complex molecules, like alcohols, into gas phase. Recently, Bottinelli et al. (2004a,b, 2006) and Kuan et al. (2004) detected relatively large abundances of complex organic molecules, like HCOOCH_3 and $(\text{CH}_3)_2\text{O}$, toward the so-called “hot corinos” associated with warm condensation surrounding low mass proto-stars.

In the following sections, we compare the abundances of complex organic molecules in the GC with those found in the hot cores and hot corinos associated with star formation.

5.1. Hot cores

For the comparison between the GC and hot cores associated to massive star formation, we selected the homogeneous data set by Ikeda et al. (2001) for several hot cores (W51, NGC6334f, G327.3–0.6, G31.41+0.31, G34.3+0.2, G10.47+0.03, and the Orion Hot Core). We also used data from

Table 5. Hot core abundances.

Source	$\bar{X}_{\text{CH}_3\text{OH}}$ $\times 10^{-7}$	$\bar{X}_{\text{C}_2\text{H}_5\text{OH}}$ $\times 10^{-8}$	$\bar{X}_{(\text{CH}_3)_2\text{O}}$ $\times 10^{-8}$	$\bar{X}_{\text{HCOOCH}_3}$ $\times 10^{-8}$	\bar{X}_{HCOOH} $\times 10^{-9}$	$\bar{X}_{\text{CH}_3\text{COOH}}$ $\times 10^{-9}$	$\bar{X}_{\text{H}_2\text{CO}}$ $\times 10^{-8}$	$N(\text{H}_2)$ $\times 10^{22} \text{ cm}^{-2}$
High mass ^a								
W51 e1/e2	3.0	0.9		3.0		0.3–1.8		36
NGC6334f	2.0	0.9	4.0		1.0			20
G327.3–0.6	1.0	1.0	3.0		0.9			20
G31.41+0.31	0.9	2.0	2.0		1.0			16
G34.3+0.2	0.9	0.6	1.0	0.5		0.2		30
G10.47+0.03	2.0	1.0	3.0		1.0			13
Omc HC	10.0	2.0		9.0				10
Low mass ^b								
IRAS 4A	≤ 0.1		≤ 2.8	6.8	4.6		2.0	160
IRAS 4B	7.0		≤ 120.0	220.0	≤ 1000		300.0	8.1
IRAS 2A	3.0		3.0	≤ 134.0	≤ 120		20.0	21
IRAS 16293	1.0		24.0	34.0	~ 62		10.0	7.5

^a Ikeda et al. (2001) and Remijan et al. (2002, 2003); ^b Bottinelli et al. (2006).

Table 6. Summary of chemical models and observations of complex organic molecules.

Source	T (K)	$n(\text{H}_2)$ (cm^{-3})	$\frac{\text{C}_2\text{H}_5\text{OH}}{\text{CH}_3\text{OH}}$ ^a $\times 10^{-2}$	$\frac{(\text{CH}_3)_2\text{O}}{\text{CH}_3\text{OH}}$ ^a $\times 10^{-2}$	$\frac{\text{HCOOCH}_3}{\text{CH}_3\text{OH}}$ ^a $\times 10^{-2}$	$\frac{\text{H}_2\text{CO}}{\text{CH}_3\text{OH}}$ ^a	age $\times 10^5 \text{ y}$
Galactic center clouds	~ 100 ^b	$\sim 10^4$ – 10^5	~ 3.66	~ 3.87	~ 3.96	$\sim 1.10 \times 10^{-2}$	
hot cores ^c	~ 100	$\sim 10^6$	~ 6.90	~ 21.80	~ 8.03		
Leung et al. (1984) ^d	10	10^4	0.14	6.00		43.6	3.16
Herbst & Leung (1986) ^e	10	10^4	2.18	5.25		9.75	3.20
Herbst & Leung (1989) ^e	10	10^4	0.99	0.12		65.5	1.00
Millar et al. (1991) ^d	70	2×10^5	0.19	6.32×10^{-2}	1.26	1.18×10^3	1.60
Hasegawa & Herbst (1993) ^f	10	10^4	0.77	1.56×10^{-2}		4.06×10^2	3.20
Caselli et al. (1993)-CR ^g	100	10^6	1.5×10^{-4}	0.21	4.67×10^{-3}	1.05	1.00
Charnley et al. (1995)	100	2×10^6		~ 2	~ 1.5	4×10^{-2}	~ 0.30
Rodgers & Charnley (2001) ^h	100	10^7	3.75	30.00	10.00		0.12
Rodgers & Charnley (2001) ⁱ	100	10^7	20.00	12.50	0.50		0.60
Horn et al. (2004) ^j	100	10^6		0.75	3.33×10^{-2}	1.56	0.80
Peeters et al. (2006) ^k	100	10^7		50.00			0.20

^a The relative abundances of $\text{C}_2\text{H}_5\text{OH}$, $(\text{CH}_3)_2\text{O}$, HCOOCH_3 , and H_2CO with respect to CH_3OH predicted by time-dependent chemical models, refer to the age of maximum abundance of the CH_3OH 's daughters; ^b from Hüttemeister et al. (1993) and Rodríguez-Fernández et al. (2001a); ^c mean abundance ratios for the hot core sources used in this work; ^d model 1 of the referred paper; ^e low-metal model in the referred paper; ^f model starting with all the H in H_2 ; ^g model fixed with the Orion Compact ridge observations; ^h model without NH_3 ; ⁱ model with NH_3 ; ^j model 3 of the referred paper; ^k model with $(\text{CH}_3)_2\text{O}$ created only in gas phase.

Remijan et al. (2002, 2003) to obtain the CH_3COOH abundances derived from their $[\text{HCOOCH}_3/\text{CH}_3\text{COOH}]$ ratio. The hot cores abundances of complex molecules and the H_2 column densities are given in Table 5. Both the H_2 and molecular column densities were derived from observations with similar beam sizes. Then the derived fractional abundances are independent of beam dilution. Since our comparison is based on the relative abundance between complex molecules, our results are not affected by the size of the hot cores.

We also used our data for the Sgr B2N and Sgr B2M hot cores in Table 4. Figure 8 illustrates the comparison between the abundance ratios observed for the GC and those measured in hot cores. As for the GC clouds, the abundance of CH_3OH in hot cores shows variations of more than one order of magnitude between different objects. Surprisingly, the abundance ratios relative to CH_3OH are also independent of the CH_3OH abundance and similar to those measured for the GC clouds. The $\text{C}_2\text{H}_5\text{OH}$ and HCOOH mean abundance ratios with respect to CH_3OH between hot cores and GC clouds are different by less than a factor of 2. The molecule that presents the largest difference in its

abundance ratio with respect to that in the GC clouds is $(\text{CH}_3)_2\text{O}$ with a mean ratio of 2×10^{-1} , which is ~ 6 times higher than those of the GC clouds. The $[\text{CH}_3\text{COOH}/\text{CH}_3\text{OH}]$ ratios in hot cores agree with the upper-limit ratios found in GC clouds. Our data for the Sgr B2N Hot Core shows slightly higher abundances of HCOOCH_3 and HCOOH than the rest of the hot cores. The Sgr B2M Hot Core, with the lowest CH_3OH abundance, shows similar ratios to those in the GC sources.

It is remarkable that different objects like the hot cores and the GC clouds, with different properties in terms of densities, H_2 column densities, and spatial extension, show such uniform abundance ratios of very complex molecules. Hot cores are denser than the GC clouds by more than 2 orders of magnitude, and they typically have larger column densities than the GC clouds.

5.2. Hot corinos

Bottinelli et al. (2004a,b, 2006), Jørgensen et al. (2005) and Kuan et al. (2004) observed the hot corinos associated with the

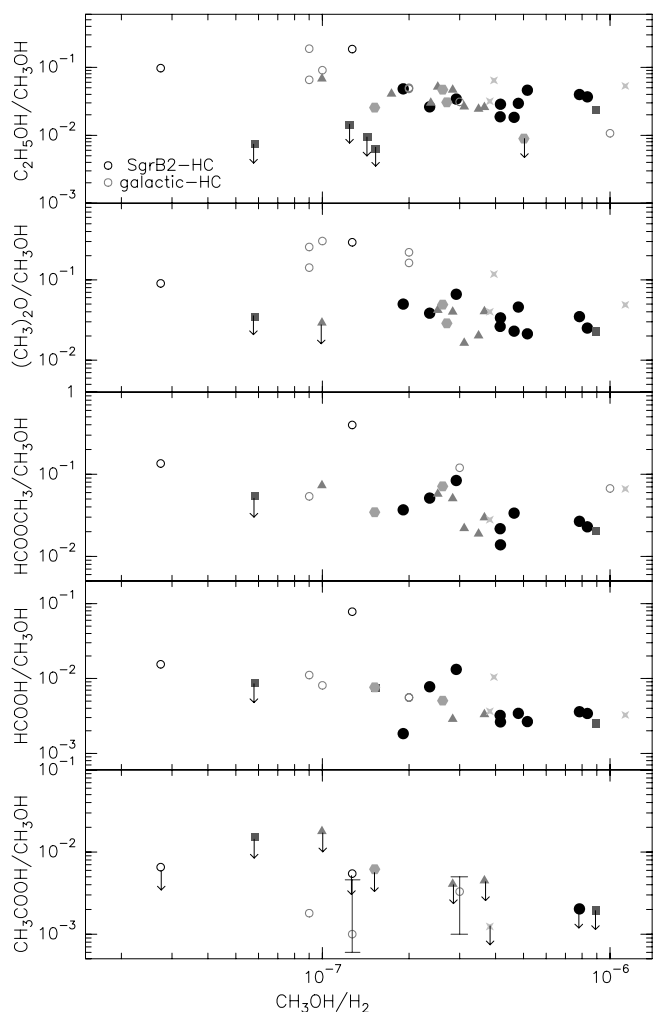


Fig. 8. Same as Fig. 5 but including the hot core abundances obtained from Ikeda et al. (2001), Remijan et al. (2002, 2003), and from our SGR B2 data as open symbols.

low mass proto-stars, NGC1333 IRAS 4A, NGC1333 IRAS 4B, NGC1333 IRAS 2A and IRAS 16293-2422 (A and B), which also show large abundances of HCOOCH₃, H₂CO, (CH₃)₂O and HCOOH. Table 5 summarizes the recent compilation of the hot corino abundances of complex organic molecules by Bottinelli et al. (2006). In this case the molecular abundances were derived by assuming a source size of 0.5'' for IRAS 4A (Bottinelli et al. 2004a), 0.25'' for IRAS 4B, 0.43'' for IRAS 2A (Maret et al. 2004), and 2'' for IRAS 16293-2422 (Cazaux et al. 2003).

Except for IRAS 4A, which shows lower abundances, HCOOCH₃, (CH₃)₂O, HCOOH, and H₂CO show higher abundances in hot corinos than in the GC molecular clouds and hot cores. The data show that the abundance of CH₃OH varies between sources by nearly two orders of magnitude, similar to the variation observed in the GC clouds. As in the GC molecular clouds and in the hot cores, the abundance of the complex organic molecules relative to that of CH₃OH is independent of the CH₃OH abundance for different hot corinos (Bottinelli et al. 2006).

Although the number of observations of hot corinos is still limited, it seems that the abundance ratios, including species believed to be formed by different chemistries, show a similar behavior to the GC sources. However, the abundance of the molecules is very different, as expected if the grain mantle

composition in dark clouds were different than in the GC clouds and hot cores.

6. Discussion

With the exception of the clouds associated with the Sickle and the TRA, large abundances of organic complex molecules like CH₃OH, C₂H₅OH, (CH₃)₂O, HCOOCH₃, HCOOH, and H₂CO seems to be a general characteristic of molecular clouds in the GC at scales of a few hundred parsecs. The main characteristic of the GC molecular gas is that the complex organic molecules show a constant relative abundance with respect to CH₃OH, within a factor 4–8 over the whole range of the CH₃OH abundances in the observed GC clouds. This contrasts with the [CS/CH₃OH] abundance ratio that increases by a factor of 60 when the CH₃OH abundance decreases.

We now discuss the origin of the chemistry of the complex molecules found in the GC sources, including the low [C₂H₅OH/CH₃OH] abundance ratio measured in the Sickle and the TRA clouds.

6.1. Models for gas-phase formations of organic molecules

Table 6 shows a compilation of the models proposed to explain the formation of complex organic molecules. Since gas-phase chemistry cannot account for the large abundance of alcohols, such as CH₃OH and C₂H₅OH, or for other organic molecules like the H₂CO measured in hot cores, evaporation from grain mantles has been proposed as the main formation mechanism. Most of these models assume that (CH₃)₂O and HCOOCH₃ were produced in gas phase after the ejection of CH₃OH. However, it has been recently claimed that HCOOCH₃ cannot be produced efficiently in gas phase from CH₃OH (Horn et al. 2004). The large abundances of this molecule found in the GC clouds clearly indicate that HCOOCH₃ is hardly likely to be produced in gas phase. However, the most recent model from Peeters et al. (2006) studied the possibilities for the formation of (CH₃)₂O on grain mantles and concluded that the gas-phase reactions are the dominant way to form (CH₃)₂O, while grain mantle reactions are the minor source.

From the theoretical point of view, so far, only (CH₃)₂O seems to be produced in gas phase from reactions involving CH₃OH on relatively short time scales. Assuming that the gas phase models for the (CH₃)₂O formation are correct (see references in Table 6), one would expect large changes in the relative abundances between the daughter ((CH₃)₂O) and parent (CH₃OH) molecules as a function of time after ejection/evaporation. One expects low [(CH₃)₂O/CH₃OH] abundance ratio in early times just when large abundances of CH₃OH are ejected into the gas phase and the gas phase processing starts. Low [(CH₃)₂O/CH₃OH] ratios are also expected in late times in the low CH₃OH abundance regime when an important fraction of CH₃OH has been converted into daughter molecules and both molecules have been destroyed by gas-phase reactions. Between these two regimes, one would expect the [(CH₃)₂O/CH₃OH] ratio to reach its maximum values, because the daughter molecules reach their maximum abundance.

From the hot core models, one expects variations in [(CH₃)₂O/CH₃OH] abundance ratio up to three orders of magnitude as a function of time. This contrasts with our data for the GC clouds and the hot cores, which show variations of less than one order of magnitude. Assuming that the chemical models for hot

cores are applicable in the GC, the only possibility of explaining the high abundances of complex molecules and the rather constant $[(\text{CH}_3)_2\text{O}/\text{CH}_3\text{OH}]$ abundance ratios is to consider that all GC clouds in our study, distributed over 200 pc, have undergone the ejection of CH_3OH from grain mantles nearly simultaneously, $\sim 10^5$ years ago. This possibility seems very unlikely, suggesting that $(\text{CH}_3)_2\text{O}$ is also ejected from grain mantles. Since $(\text{CH}_3)_2\text{O}$ shows similar abundances with respect to CH_3OH than the other molecules that can only be formed on grain mantles like $\text{C}_2\text{H}_5\text{OH}$ and HCOOCH_3 , our data support the scenario in which all complex organic molecules in the GC clouds, the hot cores, and also in hot corinos have been ejected/evaporated from grains.

Previously we also found differences in the behavior of the relative abundances of CS and the complex organic molecules. This behavior is supported by the data in shock regions associated with outflows, as in L1448 (Jiménez-Serra et al. 2005) and L1157 (Bachiller & Pérez Gutiérrez 1997). The CS abundance is enhanced by the shock, but the enhancement of CH_3OH is ~ 1 order of magnitude larger than that of CS. This effect of differential ejection of the molecules can explain the variation in the complex organic molecules abundances in gas phase, while the CS abundance remains marginally affected.

6.2. Grain mantle erosion by shocks and time scales for depletion

In the GC clouds, the typical dust-grain temperatures are low ($T \sim 10\text{--}30$ K, Rodríguez-Fernández et al. 2004) and the ejection of molecules to gas phase cannot be produced by evaporation like in hot cores, but by shocks with moderate velocities. The shock heats the gas and produces the sputtering of molecules on the grain mantles. For shock velocities of ~ 20 km s^{-1} , the gas temperature can reach ~ 1000 K (Kaufman & Neufeld 1996). After the shock passage, the gas rapidly cools on time scales of $\sim 10^4$ years for densities of 10^5 cm^{-3} . The complex molecules ejected from the grains will again stick to the grains or grain mantles. The time scales will depend on the sticking coefficient. A sticking coefficient of ~ 1 is expected for mean velocities corresponding to the thermal velocity at temperatures of $\lesssim 200$ K because the complex molecules will be absorbed after every collision with a grain (Tielens & Hagen 1982). However, the sticking coefficient decreases when increasing the relative velocity between the grains and the molecules (Leitch-Devlin & Williams 1985). Buch & Zhang (1991) and Masuda et al. (1998) estimated that the sticking coefficient of H is a function of the mean velocity.

The key parameter is the adsorption energy of the complex molecules, which will depend on the surface of the dust grains. Since the CH_3OH abundance in gas phase is relatively large, grain mantles are completely eroded and the most likely grain surfaces in the GC clouds would be graphite and silicates. The absorption energy of CH_3OH on carbon basal planes is 1600 K (Aikawa et al. 1996) and on silicates is 2065 K (Allen & Robinson 1977). For the typical mean velocities in the GC of $2\text{--}4$ km s^{-1} (see next section) the sticking coefficient will decrease and the time scale for sticking would be $\sim 10^5$ years on silicates and graphites. Similar time scales are found for the depletion of $\text{C}_2\text{H}_5\text{OH}$, HCOOCH_3 , and $(\text{CH}_3)_2\text{O}$ molecules with absorption energies ~ 3000 K on silicates (Allen & Robinson 1977). Short time scales for depletion, like those expected in the GC, would require a mechanism that continuously ejects molecules from the grain mantles.

6.3. Origin of the shocks in the GC

Rodríguez-Fernández et al. (2001a, 2004) have summarized the possible mechanisms for the heating and the chemistry of the molecular gas in the GC. Turbulence and low-velocity shocks are proposed as the most likely mechanism for driving the chemistry. One of the key parameters for constraining the origin of the shocks is the velocity dispersion in the GC molecular clouds. High angular-resolution observation of the envelope of Sgr B2 indicates that the warm molecular gas is highly turbulent with linewidths of ~ 4 km s^{-1} (Martín-Pintado et al. 1999). Smith et al. (2000) show that MHD turbulence creates a wide range of shock velocities, but the larger amount of them should be produced with Mach numbers between $2\text{--}4$. In the GC regions the sound speed is ~ 1 km s^{-1} (at 100 K), and then the more abundant shocks associated with turbulence should have velocities between 2 and 4 km s^{-1} , in agreement with the linewidth measured from high angular-resolution observations. As discussed in the previous section, for a mean velocity of ~ 3 km s^{-1} , the depletion will be $\sim 10^5$ years.

The energy of any kind of turbulence will decay if there is not any source injecting new energy. It is interesting to compare the depletion time scales with the turbulence decay time scales to establish if the turbulence can maintain mean velocities that are high enough to match the depletion time scales. The turbulence decays as $t^{-\alpha}$, where $\alpha \sim 0.8\text{--}1.0$ (Mac Low et al. 1998; Stone et al. 1998). The decay time scale of these turbulences is similar to one dynamical time ($t_{\text{dyn}} = L/\sigma_v$, where L is the turbulent length and σ_v the turbulent velocity, Avila-Reese & Vázquez-Semadeni 2001; Elmegreen & Scalo 2004), for both HD and MHD turbulence. The dynamical time for turbulence in the GC, assuming a turbulent length of 5 pc and a turbulent velocity dispersion of 15 km s^{-1} (observed GC linewidths), is $\sim 3 \times 10^5$ years, close to the depletion time scale. Since we do not observe large depletions in our data, the existence of faster shocks with velocities high enough to produce the sputtering of the grain mantles and with the time scales between shocks similar to the time scales for depletion, $\sim 10^5$ years is required. Shocks with velocities of ≥ 6 km s^{-1} are required in order to sputter molecules from the icy mantles on graphite and silicate grains by heavy atoms (Draine & Salpeter 1979). Then, if due to a sudden event, the turbulence generated by this event could explain the presence of complex organic molecules in the GC clouds for $\sim 10^5$ years.

It is so far unclear what produces the supersonic turbulence and/or frequent shocks in the GC. The high-velocity shocks and the presence of a highly turbulent medium in the GC could be due to: a) the kinematics of the gas subjected to the barred potential of our Galaxy, b) the wind-blown bubbles produced in evolved massive stars like supernovas, or c) cloud-cloud collisions. The time scales needed in the GC to maintain the large abundances are much smaller than those associated to the quasi-circular orbits ($\sim 1 \times 10^7$ years), and a large number of large scale shocks would be required to explain the observed properties of the complex molecules. The other possible mechanisms could explain the observations if cloud-cloud collision and energetic events driven by massive stars occur on time scales of 10^5 years. In the scenario of a recent star burst in the GC as proposed to explain the fine structure lines of ionized gas (Rodríguez-Fernández & Martín-Pintado 2005), frequent energetic events associated to massive stars are expected to produce shocks with moderate velocities of ~ 10 km s^{-1} (Martín-Pintado et al. 1999) ejecting complex molecules to gas phase and a large amount of energy into the ISM.

6.4. The chemistry of complex molecules in the Galaxy

The proposed gas-phase chemistry for some complex molecules does not seem to explain the similar abundances ratios observed in the GC clouds, in hot cores, and in hot corinos. From the empirical data gathered there, we can conclude that the most likely explanation for the large abundance of CH_3OH , HCOOH , HCOOCH_3 , $(\text{CH}_3)_2\text{O}$ and $\text{C}_2\text{H}_5\text{OH}$ is that all these molecules are ejected from grains. It is interesting to note that in this scenario, the ice-mantle composition in complex organic molecules in the GC and in hot cores must be very similar. This is surprising in view of the changes of the complex organic molecules abundances observed in the hot corinos. The presence of complex molecules on grains can be due to grain chemistry or to the depletion after formation in gas phase.

Further investigations of the abundance of organic molecules in dark clouds are needed to constrain the molecules formed on ice mantles by grain chemistry in different types of molecular clouds.

6.5. Sources with low $\text{C}_2\text{H}_5\text{OH}$ abundances

We also observed some clouds where the relative abundance of some organic complex molecules like $\text{C}_2\text{H}_5\text{OH}$ seems to be significantly smaller than expected from these observed in other sources in the GC. These sources are the Sickie (MC G+0.18+0.04 and MC G+0.20+0.03) and the -30 km s^{-1} velocity component of the TRA (MC G+0.13+0.02). We detected high CH_3OH abundances of $\sim 10^{-7}$, but only significant upper limits, $\lesssim 0.4\text{--}5 \times 10^{-9}$ to the $\text{C}_2\text{H}_5\text{OH}$ abundance. The stars from the Quintuplet and the Arches Clusters are said to heat and ionize the Sickie and the TRA (Rodríguez-Fernández et al. 2001b) creating photodissociation-regions (PDRs) around them. In particular, the -30 km s^{-1} component is clearly associated with the TRA (Serabyn & Güsten 1987).

The chemistry in these sources is then expected to be strongly affected by the presence of the UV radiation. It is very likely that the complex molecules in these sources are ejected to gas phase by shocks with the same abundances as observed in other GC molecular cloud. Additional evaporation from grain mantles can be ruled out since the heating will require UV photons that will also destroy the complex molecules. Due to a large UV radiation-field, the relative abundance of the complex molecules will be affected by dissociation. One can make a rough estimate of the effects of the UV radiation on the molecular abundances by comparing their photodissociation rates. Using the UMIST RATE99 (Le Teuff et al. 2000) and the Ohio State University databases¹, we found that the photodissociation rates of $(\text{CH}_3)_2\text{O}$ and $\text{C}_2\text{H}_5\text{OH}$ are larger than those of CH_3OH , HCOOH and H_2CO .

In Fig. 9 we show the results of a simple model in which we represent how the ejected complex organic molecules are photodestroyed as a function of time and visual extinction. The model considers a far-ultraviolet incident field 10^3 times higher than in the local ISM ($G_0 = 10^3$), which is the typical radiation field observed in GC PDRs (Rodríguez-Fernández et al. 2004). For the regions with low A_v , all the abundances decrease, but the abundance of $\text{C}_2\text{H}_5\text{OH}$ decreases faster than the other organic complex molecules. Obviously the time scale for photodissociation depends on the visual extinction. We estimate a visual extinction of $A_v \sim 4\text{--}17$ mag for the Sickie and the TRA using our H_2 column densities in Table 4 and the conversion factor given

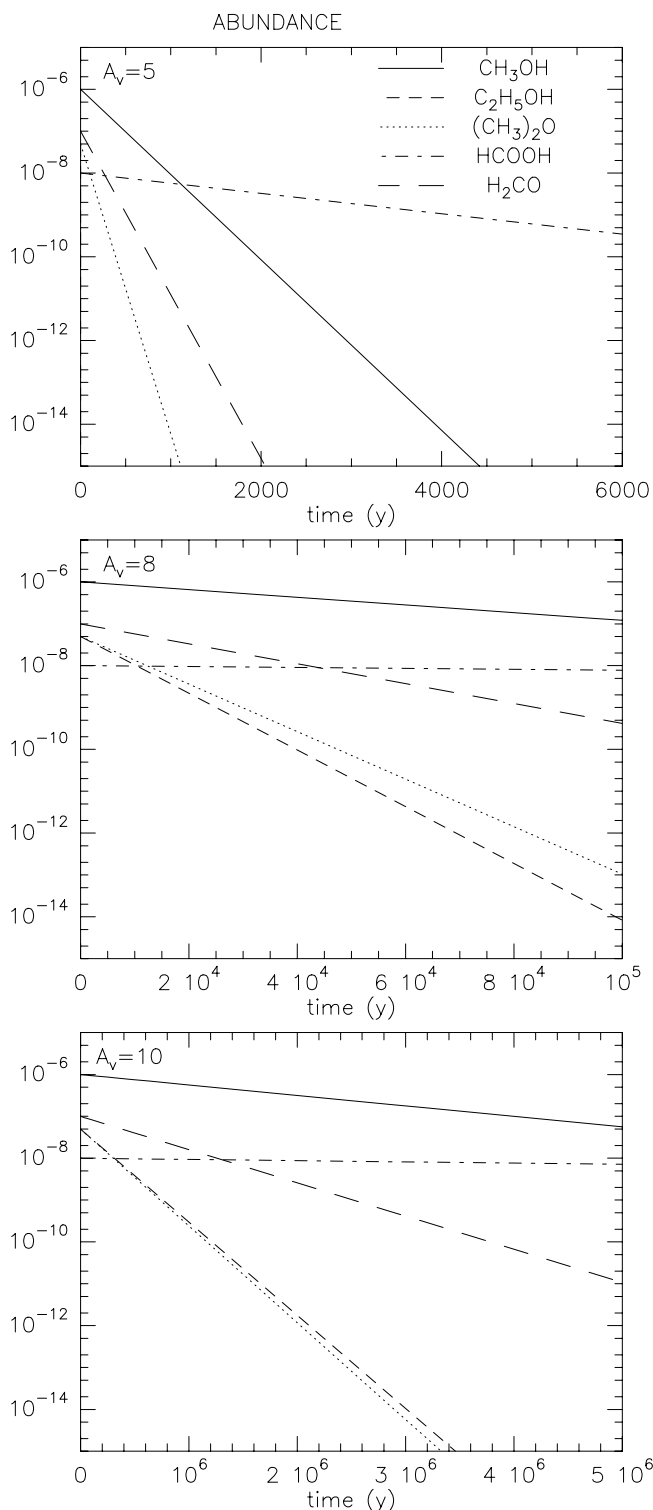


Fig. 9. Photodestruction of different complex molecules as a function of time for different visual extinctions (A_v). The initial molecular abundances for the molecules are considered to be those typically measured in the observed GC clouds. $G_0 = 10^3$, observed by Rodríguez-Fernández et al. (2004), is used for the UV radiation field.

by Bohlin (1975), $A_v(\text{mag}) = 10^{-21} \times N_{\text{H}_2} (\text{cm}^{-2})$. The photodissociation time scales for CH_3OH will be comparable to depletion time scales for an averaged visual extinction of ~ 8 mag.

For this visual extinction the other complex organic molecules will photodissociate much faster than the depletion

¹ www.physics.ohio-state.edu/~eric/research.html

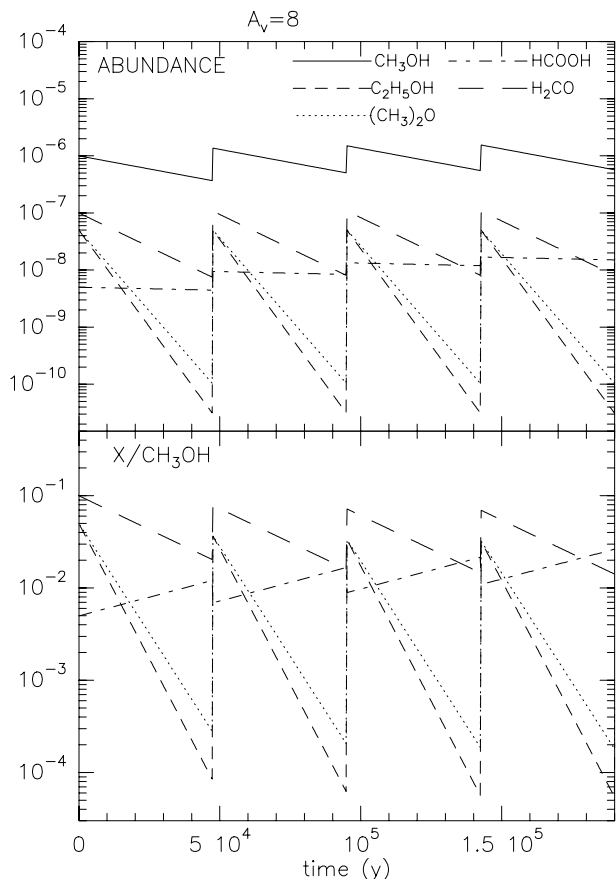


Fig. 10. Time evolution of the abundance (*top*) and relative abundance (*bottom*) of complex organic molecules with respect to that of CH_3OH in a PDR with $A_v = 8$ mag where a periodic shock every 5×10^4 years enhances the abundances to their initial values. The radiation field is $G_0 = 10^3$.

time scale. This suggests that in the presence of UV radiation, large abundances of molecules like $\text{C}_2\text{H}_5\text{OH}$ would require more frequent shocks than in the shielded regions. This is illustrated in Fig. 10, where we present the results of a simple model to show the time evolution of complex organic molecules in a region of $A_v = 8$ mag. In this model, we consider the effects of the UV radiation on the complex molecules that have been ejected by periodic shocks of 5×10^4 years. They are ejected with the abundances measured in the GC and are destroyed because of the photodissociation. This simple model reproduces the constant high abundances observed for CH_3OH , while the abundances of $\text{C}_2\text{H}_5\text{OH}$ can vary by ~ 3 orders of magnitude. In agreement with the observations, the photodissociation can reproduce the low abundances of $\text{C}_2\text{H}_5\text{OH}$ in specific regions in the GC clouds. Further observation of complex organic molecules toward PDRs in the Galactic disk and in the Galactic center are required to test the proposed scenario.

7. Conclusion

We have found very high abundances for the complex organic molecules CH_3OH , $\text{C}_2\text{H}_5\text{OH}$, HCOOCH_3 , $(\text{CH}_3)_2\text{O}$, and HCOOH in the GC regions, similar to or even higher than those measured in Galactic hot cores. The abundance ratios of these molecules are fairly constant in all the GC clouds. The most likely explanation for these large abundances is that they have been ejected from grain mantles by shocks.

The highly turbulent molecular clouds in the GC could help to maintain the large gas phase abundance of complex molecules for time scales of $\sim 10^5$ years after the ejection by shocks. The rather uniform abundance ratios in clouds distributed over 200 pc indicate that the average composition of grain mantles is similar for all the GC clouds and also for the hot cores in the Galactic disk. We found that, in the GC PDRs, UV photons can differentially photodissociate the complex molecules, changing their relative abundances dramatically.

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