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Methanol deuteration in high-mass protostars

M. L. van Gelder¹, J. Jaspers¹, P. Nazari¹, A. Ahmadi¹, E. F. van Dishoeck^{1,2}, M. T. Beltrán³, G. A. Fuller^{4,5}, Á. Sánchez-Monge⁵, and P. Schilke⁵

- ¹ Leiden Observatory, Leiden University, PO Box 9513, 2300RA Leiden, The Netherlands e-mail: vgelder@strw.leidenuniv.nl
- ² Max Planck Institut für Extraterrestrische Physik (MPE), Giessenbachstrasse 1, 85748 Garching, Germany
- ³ INAF-Osservatorio Astrofisico di Arcetri, Largo E. Fermi 5, 50125 Firenze, Italy
- ⁴ Jodrell Bank Centre for Astrophysics, Department of Physics and Astronomy, University of Manchester, Oxford Road, Manchester M13 9PL, UK
- ⁵ I. Physikalisches Institut, Universität zu Köln, Zülpicher Str.77, 50937 Köln, Germany

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ABSTRACT

Context. The deuteration of molecules forming in the ices such as methanol (CH₃OH) is sensitive to the physical conditions during their formation in dense cold clouds and can be probed through observations of deuterated methanol in hot cores.

Aims. The aim is to determine the D/H ratio of methanol for a large sample of 99 high-mass protostars and to link this to the physical conditions during the formation of methanol in the prestellar phases.

Methods. Observations with the Atacama Large Millimeter/submillimeter Array (ALMA) containing transitions of CH₃OH, CH₂DOH, CHD₂OH, ¹³CH₃OH, and CH₃¹⁸OH are investigated. The column densities of CH₂DOH, CHD₂OH, and CH₃OH are determined for all sources, where the column density of CH₃OH is derived from optically thin ¹³C and ¹⁸O isotopologues. Consequently, the D/H ratio of methanol is derived taking statistical effects into account.

Results. Singly deuterated methanol (CH₂DOH) is detected at the 3σ level toward 25 of the 99 sources in our sample of the high-mass protostars. Including upper limits, the (D/H)_{CH₃OH} ratio inferred from $N_{\text{CH}_2\text{DOH}}/N_{\text{CH}_3\text{OH}}$ was derived for 38 of the 99 sources and varies between $\sim 10^{-3} - 10^{-2}$. Including other high-mass hot cores from the literature, the mean methanol D/H ratio is $1.1 \pm 0.7 \times 10^{-3}$. This is more than one order of magnitude lower than what is seen for low-mass protostellar systems ($2.2 \pm 1.2 \times 10^{-2}$). Doubly deuterated methanol (CHD₂OH) is detected at the 3σ level toward 11 of the 99 sources. Including upper limits for 15 sources, the (D/H)_{CH₂DOH} ratios derived from $N_{\text{CHD}_2\text{OH}}/N_{\text{CH}_2\text{DOH}}$ are more than two orders of magnitude higher than (D/H)_{CH₃OH} with an average of $2.0 \pm 0.8 \times 10^{-1}$ which is similar to what is found for low-mass sources. Comparison with literature GRAINOBLE models suggests that the high-mass prestellar phases are either warm (>20 K) or live shorter than the free-fall timescale. In contrast, for low-mass protostars, both a low temperature of <15 K and a prestellar phase timescale longer than the free-fall timescale are necessary.

Conclusions. The $(D/H)_{CH_3OH}$ ratio drops by more than an order of magnitude between low-mass and high-mass protostars due to either a higher temperature during the prestellar phases or shorter prestellar phases. However, successive deuteration toward CHD_2OH seems equally effective between low-mass and high-mass systems.

Key words. astrochemistry – stars: formation – stars: protostars – techniques: interferometric – ISM: abundances

1. Introduction

Isotopologues have proven to be vital in our understanding of the star and planet formation process. They allow for the most abundant species to be studied for which the emission originating from the main isotopologue is optically thick. Moreover, the sensitivity of isotopologue ratios to the physical conditions such as temperature and ultraviolet (UV) radiation has proven key in understanding the molecular journey during the entire star formation process (see e.g., reviews by Caselli & Ceccarelli 2012; Tielens 2013; Ceccarelli et al. 2014). One of the most studied types of isotopologues are those that contain deuterium (D). These deuterated molecules are suggested to form already in the cold prestellar phases (e.g., van Dishoeck et al. 1995; Caselli & Ceccarelli 2012; Ceccarelli et al. 2014). Especially for molecules such as methanol (CH₃OH) that form on the surfaces of dust grains in dense cores, D/H fractionation ratios up to 10% are found toward low-mass protostars (e.g., Bianchi et al. 2017a,b; Taquet et al. 2019; van Gelder et al. 2020), more than four orders of magnitude larger than the canonical D/H ratio derived for the

local interstellar medium (ISM) of of $\sim 2 \times 10^{-5}$ (Linsky et al. 2006; Prodanović et al. 2010). It is thus key to understand the deuterium fractionation process in the earliest phases of star formation.

The gaseous atomic D/H ratio can be increased in the prestellar phases through the exothermic reaction (Watson 1974; Aikawa & Herbst 1999; Ceccarelli et al. 2014),

$$H_3^+ + HD \rightleftharpoons H_2D^+ + H_2 + \Delta E, \tag{1}$$

where $\Delta E = 232$ K. Since in the cold ($\lesssim 20$ K) prestellar cores the backward reaction in Eq. (1) is less efficient, H_2D^+ is enhanced and the atomic D/H ratio in the gas phase can be effectively increased through dissociative recombination of H_2D^+ with free electrons. Moreover, gaseous CO is the main destructor of H_3^+ and H_2D^+ (Brown & Millar 1989; Roberts et al. 2003) and thus the heavy CO freeze-out in dense ($\gtrsim 10^4$ cm⁻³) prestellar cores additionally stimulates the increase of the gaseous atomic D/H ratio. In turn, the enhanced atomic D/H ratio in the gas can translate into a higher D/H ratio of molecules forming in the

ices (Tielens 1983; Nagaoka et al. 2005). Measuring the deuteration of molecules that form in the ices is thus a powerful tool to determine the physical conditions such as density (e.g., CO freeze-out) and temperature during their formation.

Methanol forms on the surfaces of dust grains in dense prestellar phases through the hydrogenation of CO ice (e.g., Watanabe & Kouchi 2002; Fuchs et al. 2009) and reactions between its grains-surface products (e.g., H₂CO and CH₃O; Simons et al. 2020; Santos et al. 2022) and is therefore expected to exhibit a high D/H ratio. This is in strong contrast to, for example, water for which the bulk of the ice is formed in the warmer translucent cloud phase leading to a rather low overall HDO/ H_2O ratio ($\lesssim 0.1\%$; Persson et al. 2014; Furuya et al. 2016; Jensen et al. 2019; van't Hoff et al. 2022). The sensitivity of the methanol deuteration process to temperature was investigated by Bøgelund et al. (2018) using the GRAINOBLE gas-grain chemical model (Taquet et al. 2012, 2013, 2014), finding a strong correlation between the D/H ratio of methanol and the formation temperature. Moreover, Taquet et al. (2019) showed that the timescale of the prestellar phase is highly relevant for methanol deuteration.

Methanol and its (deuterated) isotopologues are readily observed as they desorb from the dust grains. Mono deuterated methanol, CH2DOH and CH3OD, have been observed in the warm inner regions of both low-mass and high-mass protostellar systems (e.g., Fuente et al. 2014; Belloche et al. 2016; Bøgelund et al. 2018; van Gelder et al. 2020; van der Walt et al. 2021). Similarly, both doubly and triply deuterated methanol have been detected in hot cores (e.g., Parise et al. 2002, 2004; Bianchi et al. 2017a; Drozdovskaya et al. 2022; Ilyushin et al. 2022). Moreover, CH₂DOH has also been detected in both lowmass prestellar cores (e.g., Bizzocchi et al. 2014; Lattanzi et al. 2020; Ambrose et al. 2021) and high-mass starless cores (e.g., Fontani et al. 2015). Across this mass and evolutionary range, the D/H ratio of singly deuterated methanol varies orders of magnitude. The D/H ratio is on the order of 10% for low-mass prestellar cores, low-mass protostars, and comets (e.g., Bianchi et al. 2017a,b; Jørgensen et al. 2018; Taquet et al. 2019; Manigand et al. 2020; van Gelder et al. 2020; Lattanzi et al. 2020; Ambrose et al. 2021; Drozdovskaya et al. 2021). Interestingly, successive deuteration toward CHD₂OH and CD₃OH seems to be quite effective in low-mass protostars (about 15–25%; Drozdovskaya et al. 2022; Ilyushin et al. 2022). On the other hand, the D/H ratio derived from CH₂DOH is as low as 0.1-0.01% for high-mass starless cores and high-mass protostars (Fontani et al. 2015; Neill et al. 2013; Belloche et al. 2016; Bøgelund et al. 2018). However, the sample of high-mass protostars for which reliable and interferometrically derived methanol D/H ratios are available (e.g., Orion KL, Sgn B2(N2), NGC 6334I) remains small compared to the low-mass sources (~20 sources). Furthermore, no interferometric detections of CHD₂OH in high-mass sources have been presented thus far.

In this work, the methanol D/H ratios are derived for an additional 99 high-mass sources based on ALMA observations of CH₂DOH, CHD₂OH, CH₃OH ¹³CH₃OH, CH₃¹⁸OH. In Sect. 2, the observations and derivation of the column densities are explained. The resulting D/H ratios of CH₃OH and CH₂DOH are presented in Sect. 3. In Sect. 4, the methanol D/H ratios derived for our high-mass sources are compared to their low-mass counterparts and prestellar phases. Furthermore, through comparison with the GRAINOBLE models computed by Bøgelund et al. (2018) and Taquet et al. (2019), the effect of physical conditions on the methanol D/H ratio is discussed. Our main conclusions are listed in Sect. 5.

2. Methodology

2.1. Observations

The dataset analyzed in this work was taken from the ALMA Evolutionary study of High Mass Protocluster Formation in the Galaxy (ALMAGAL) survey (2019.1.00195.L; PI: S. Molinari) that targeted over 1000 dense clumps with $M > 500 M_{\odot}$ based on the Herschel Hi-Gal survey (Molinari et al. 2010; Elia et al. 2017, 2021). The ALMAGAL survey covers frequencies from ~217 GHz to ~221 GHz with multiple configurations of ALMA down to ~0.1" resolution at a spectral resolution of between $\sim 0.2-0.7$ km s⁻¹. In this work a subsample of 40 high-mass cores is selected based on high bolometric luminosity (L_{bol} > 1000 L_{\odot}) and the sources being rich in lines from complex organic molecules (COMs) such as CH₃OH and CH₃CN. Only archival data with a beam smaller than 2" (~1000-5000 au) that were public before February 2021 are included. This selection introduces a bias in our sample to line-rich sources and means that not all high-mass cores in the ALMAGAL survey are covered. In the higher resolution ALMA data, the 40 Hi-Gal high-mass cores are resolved into in total 99 sources based on the continuum emission (labeled A, B, C, etc., see Appendix B). These 99 sources are all studied in this work and are the same as those that were analyzed by van Gelder et al. (2022). The data were pipeline calibrated and imaged with the Common Astronomy Software Applications¹ (CASA; McMullin et al. 2007) version 5.6.1. The angular resolution of the data ranges from 0.5-1.25", corresponding to about ~2500–10000 au at the range of distances covered (2–12 kpc; Mège et al. 2021), and the data have a sensitivity of ~0.2 K. The ALMAGAL data cover several transitions of CH₃OH, four transitions of ¹³CH₃OH, nine transitions of CH₂¹⁸OH, 21 transitions of CH₂DOH, and 22 transitions of CHD₂OH (see Appendix A). Also nine transitions of CD₃OH are covered (Ilyushin et al. 2022), but these are not detected toward any of the sources. No transitions of CH₃OD are covered in the observed frequency range. Both CD₃OH and CH₃OD are therefore not analyzed further in this paper.

Integrated intensity maps of the CH₃OH 8_{0,8}-7_{1,6}, CH₂DOH $17_{1,16} e_0 - 17_{0,17} e_0$, and CHD₂OH $7_{0,1} e_1 - 6_{1,1} e_1$ lines for the source 881427 are presented in Fig. 1. The source 881427 hosts three nearby hot cores with varying line strengths and line widths and is a representative source of the rest of the sample. Whereas the emission of CH₃OH is often larger than the central beam, the emission of both CH2DOH and CHD2OH is generally confined within the central beam similar to the ^{13}C and ^{18}O isotopologues. The main exception for this is $CH_2DOH \, 5_{1,5} \, e_0 - 4_{1,4} \, e_0 \, (E_{up} = 36 \, \text{K})$ which often shows more extended emission. This is likely because the low upper energy level of this line is also sensitive to cold ($T \leq 70$ K) material where methanol is nonthermally desorbed from the grains (e.g., Perotti et al. 2020, 2021). To exclude the contribution of extended emission, this transition is not included in the analysis described below.

The spectra were extracted from the peak pixel in the CH₃OH $8_{0.8}$ – $7_{1.6}$ ($E_{\rm up}=97$ K) integrated intensity maps for all sources that show this at the $>3\sigma$ level. This line is strongest transition of CH₃OH in the sample with $E_{\rm up}>70$ K (lines with lower $E_{\rm up}$ can suffer from contamination by the outflow or extended emission). For sources that do not show emission from the CH₃OH $8_{0.8}$ – $7_{1.6}$ line, spectra were extracted from the peak continuum pixel and only upper limits on the column densities of CH₃OH (and isotopologues) are derived. In the G323.7399-00.2617B

https://casa.nrao.edu/

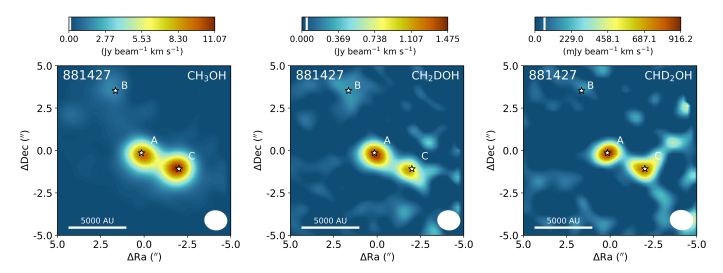


Fig. 1. Integrated intensity maps of the CH₃OH $8_{0.8}$ – $7_{1.6}$ ($E_{up} = 97$ K, left), CH₂DOH $17_{1.16}$ $e_0 - 17_{0.17}$ e_0 ($E_{up} = 336$ K, middle), and CHD₂OH $7_{0.1}$ $e_1 - 6_{1.1}$ e_1 ($E_{up} = 74$ K, right) lines for 881427. The color scale is shown on top of each image. The image is integrated over [-5, 5] km s⁻¹ with respect to the V_{lsr} of source A. The white vertical line in the colorbar indicates the 3σ threshold. The source positions based on the continuum emission are indicated with the white stars. The white ellipse in the lower right of each image depicts the beam size and in the lower left a physical scale bar is displayed.

cluster, which contains seven nearby cores, all spectra were extracted from the same positions as van Gelder et al. (2022). In G023.3891+00.1851, the emission of CH₂DOH peaks offset by about half the beam ($\sim\!0.6^{\prime\prime})$ and therefore the spectrum was extracted from the peak of CH₂DOH $17_{1,16} e_0 - 17_{0,17} e_0$ $(E_{\rm up} = 336 \text{ K})$. For all other sources, the peak in CH₃OH coincides with the peaks of CH₂DOH and CHD₂OH. For sources also included by Nazari et al. (2022), our spectral extraction locations are the same as theirs, which were extracted from the peak position of the CH₃CN 12₄–11₄ integrated intensity maps, except for 721992 and G023.3891+00.1851 where the CH₃OH $8_{0,8}$ – $7_{1,6}$ and CH₂DOH $17_{1,16} e_0 - 17_{0,17} e_0$ emission peaks offset from the CH₃CN 12₄-11₄ emission by about 1". It is important to note that these spectral extraction positions are different by up to 1" from van Gelder et al. (2022) who extracted their spectra from the peak continuum pixel for all sources. Therefore, the column densities derived in this work may deviate from theirs. The reason why our spectra are extracted from the peak pixel of CH₃OH is to have the highest signal-to-noise in methanol lines and its isotopologues the extracted spectra.

2.2. Deriving the column densities

The column densities of all methanol isotopologues were derived using the spectral analysis tool CASSIS² (Vastel et al. 2015) under the assumption of local thermodynamic equilibrium (LTE). The line lists of CH₃OH, 13 CH₃OH, and CH₃¹⁸OH were taken from the CDMS catalog³ (Müller et al. 2001, 2005; Endres et al. 2016). These entries include the first three (CH₃OH) and two (13 CH₃OH and CH₃¹⁸OH) torsional states and are based on the works of Xu et al. (2008), Xu & Lovas (1997), and Fisher et al. (2007), respectively. The difference between the statistical weight factors g_1 of 13 CH₃OH (g_1 = 1) and CH₃¹⁸OH and CH₃OH (g_1 = 4) is correctly taken into account in the CDMS database entries and therefore does not affect any column densities derived in this work. The line list of CH₂DOH was taken

from the JPL catalog⁴ (Pickett et al. 1998), where the entry is based on the work of Pearson et al. (2012). The line list of CHD₂OH was taken from Drozdovskaya et al. (2022), which is mostly based on the work of Coudert et al. (2021).

Only transitions with $E_{\rm up} \ge 50$ K were used to derive the column densities since lines with lower $E_{\rm up}$ likely include also emission from non-thermally desorbed methanol and emission possibly related to outflows. For ¹³CH₃OH, the 14_{1,13}–13_{2,12} $(E_{\rm up} = 254 \text{ K})$ transition gives the only constraint on the column density for many sources as the other transitions suffer from severe line blending. Similarly, for $CH_3^{18}OH$ only the $8_{1,8}-7_{0,7}$ $(E_{\rm up} = 86 \text{ K})$ and $14_{1,14}$ – $13_{2,12}$ ($E_{\rm up} = 239 \text{ K}$) transitions provide constraints on the column density as well as some information on the excitation temperature. Furthermore, for CH₂DOH the $5_{2,4}e_1 - 4_{1,5}e_1$ ($E_{\rm up} = 59$ K) line, as well as several other lines, have rather low Einstein A_{ij} values ($<10^{-5}~{\rm s}^{-1}$) and are often blended with other COMs. Moreover, the spectroscopy of the CH₂DOH $18_{1,17} o_1 - 18_{2,17} e_0$ line is unreliable and shows large discrepancies in A_{ij} between the JPL catalog entry (A_{ij} = 1.8×10^{-5} s⁻¹) and that derived by Coudert et al. (2014, A_{ij} = 8.9×10^{-7} s⁻¹) and is therefore also excluded from the analysis. Consequently, the $17_{1,16} e_0 - 17_{0,17} e_0$ ($E_{up} = 336 \text{ K}$) transition of CH₂DOH provided the best constraint on the column density of CH₂DOH. However, although the $5_{1,5} e_0 - 4_{1,4} e_0$ ($E_{up} = 36 \text{ K}$) transition is excluded from the fitting, it can provide information on the excitation temperature of CH₂DOH as the best-fit LTE model should not overproduce this line. Lastly, for CHD₂OH, the $7_{0,1} e_1 - 6_{1,1} e_1$ ($E_{up} = 74$ K) transition is the only detected line in our sample and therefore is the only constraint on the column density of CHD₂OH.

As a consequence of only single or a few lines being available, the excitation temperature was fixed to 150 K, which is roughly the mean temperature as measured toward other highmass hot cores (e.g., Neill et al. 2013; Belloche et al. 2016; Bøgelund et al. 2018, 2019). However, if clear anticorrelations between the best-fit LTE model and the data were present, the excitation temperature was varied by eye in steps of 25 K until

http://cassis.irap.omp.eu/

https://cdms.astro.uni-koeln.de/

⁴ https://spec.jpl.nasa.gov/

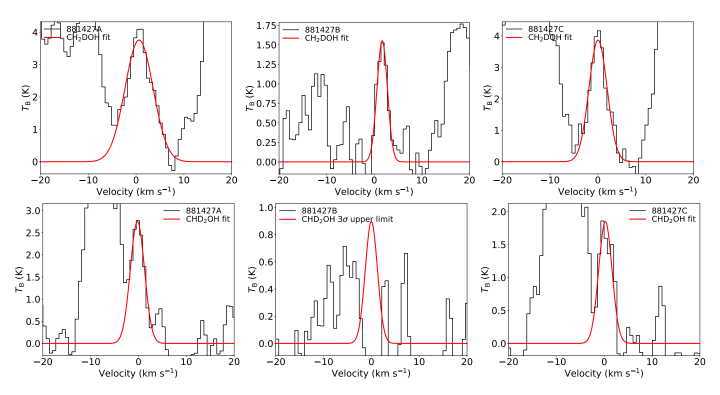


Fig. 2. Spectral line fits of CH₂DOH $17_{1,16}e_0 - 17_{0,17}e_0$ ($E_{up} = 336$ K, top row) and CHD₂OH $7_{0,1}e_1 - 6_{1,1}e_1$ ($E_{up} = 74$ K, bottom row) for 881427A (left), 881427B (middle), and 881427C (right). The data corrected for the V_{lsr} are shown in black and the best fit for $T_{ex} = 150$ K is shown in red.

the anticorrelations disappeared in a similar way as the by-eye fit method of Nazari et al. (2021, see their Appendix C).

The column densities N of ${}^{13}\text{CH}_3\text{OH}$, ${}^{18}\text{OH}$, ${}^{18}\text{OH}$, ${}^{18}\text{CH}_2\text{DOH}$, and CHD₂OH were derived following a similar method as van Gelder et al. (2020). A grid of N and the full width at half maximum (FWHM) of the line was set and a model spectrum was computed for each grid point assuming LTE conditions. The size of the emitting region was fixed to the size of the beam (see Appendix B). Blended lines were excluded from the fitting procedure and similarly broad lines (FWHM $\geq 10 \text{ km s}^{-1}$) and lines with $E_{\rm up} \leq 50$ K were excluded in the fit to exclude any emission possibly related to outflows. The best-fit column density and the 2σ uncertainty were computed from the grid for each isotopologue. The main contributors to the uncertainty of N are the uncertainty on the flux calibration of ALMA (assumed to be 10%) and the assumed excitation temperature. However, changing the excitation temperature in the 100-300 K range leads to at most a factor 3 variation in the derived column densities. For several sources (e.g., 705768), the lines are broad (>7 km $\rm s^{-1}$) making automated line fitting complicated. For these sources the column density was estimated using the by eye fitting method of Nazari et al. (2021). In this case, a 50% uncertainty on the column density was assumed. Moreover, the sources 101899, 615590, 865468, and G345.5043+00.3480 showed line profiles consisting of multiple components. The column density of each component was derived and reported separately.

For all sources, the column density of CH_3OH was derived from $\text{CH}_3^{18}\text{OH}$ and, when no lines originating from $\text{CH}_3^{18}\text{OH}$ were detected, from $^{13}\text{CH}_3\text{OH}$. The adopted $^{12}\text{C}/^{13}\text{C}$ and $^{16}\text{O}/^{18}\text{O}$ ratios are dependent on the galactocentric distance and are determined using the relations of Milam et al. (2005) and Wilson & Rood (1994), respectively. In cases where only upper limits on the column densities of both $^{13}\text{CH}_3\text{OH}$ and $\text{CH}_3^{18}\text{OH}$ could be derived, the range in $N_{\text{CH}_3\text{OH}}$ was calculated by

setting the 3σ upper limit based on scaling the 3σ upper limit of $^{13}\text{CH}_3\text{OH}$ and the lower limit based on the main isotopologue. Lastly, when CH₃OH was not detected, the 3σ upper limit was derived directly from CH₃OH lines.

3. Results

The derived column densities of all isotopologues are presented in Table B.1 for the reported excitation temperature. In Fig. 2, the best-fit models to the CH₂DOH $17_{1,16} e_0 - 17_{0,17} e_0$ and CHD₂OH $7_{0,1} e_1 - 6_{1,1} e_1$ lines are presented for three hot cores in 881427 (see Fig. 1). Toward 25 sources, at least one clean unblended line of CH₂DOH is detected at the 3σ level, allowing for the determination of the column density. For the remaining 74 sources where no (unblended) transitions of CH₂DOH are detected, the 3σ upper limit is reported. For CHD₂OH, the column density could be determined for 11 sources. In Table B.1, the column densities of 13 CH₃OH, CH $_3^{18}$ OH, and CH₃OH are also reported.

The column densities of CH_2DOH are generally between one and three orders of magnitude lower than those of CH_3OH , see also Fig. 3. Furthermore, the column densities of CHD_2OH are about a factor 3–10 lower than that of CH_2DOH , see Fig. 4. In order to translate the column density ratios to the D/H ratios, statistical weighting has to be taken into account since a deuterium atom has a three times higher probability to land in the CH_3 group compared to the OH group. Therefore, the D/H ratios of CH_3OH and CH_2DOH can be derived through,

$$N_{\text{CH}_2\text{DOH}}/N_{\text{CH}_3\text{OH}} = 3(D/H)_{\text{CH}_3\text{OH}},$$
 (2)

$$N_{\text{CHD}_2\text{OH}}/N_{\text{CH}_2\text{DOH}} = (D/H)_{\text{CH}_2\text{DOH}}.$$
 (3)

The derived D/H ratios are also listed in Table B.1.

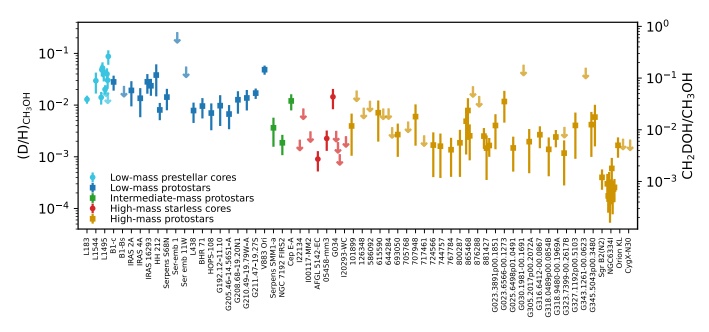


Fig. 3. $(D/H)_{CH_3OH}$ ratio (squares) derived from the N_{CH_2DOH}/N_{CH_3OH} for low-mass, intermediate-mass, and high-mass protostellar systems including data from both this study and the literature (see Appendix C for references). Only D/H ratios derived from interferometric observations are included to minimize effects of beam dilution and to exclude any contribution from larger scales. Also data for low-mass prestellar cores and high-mass starless cores (circles) from the literature (including observations with single dish telescopes) are included. Upper limits are presented as arrows.

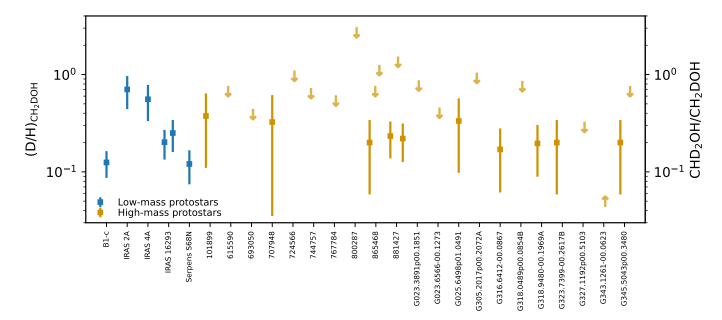


Fig. 4. $(D/H)_{CH_2DOH}$ ratio derived from the N_{CHD_2OH}/N_{CH_2DOH} for low-mass and high-mass protostellar systems including data from both this study and the literature (see Appendix C for references). Upper and lower limits are presented as arrows. Only D/H ratios derived from interferometric observations are included to minimize effects of beam dilution and to exclude any contribution from larger scales. The $(D/H)_{CH_2DOH}$ of IRAS 2A and IRAS 4A were derived using older spectroscopic data (Taquet et al. 2019).

The resulting (D/H)_{CH₃OH} and (D/H)_{CH₂DOH} ratios are presented in Figs. 3–4, respectively. Including upper limits, a (limit on the) (D/H)_{CH₃OH} and (D/H)_{CH₂DOH} ratios could be derived for 38 and 26 of the 99 studied sources, respectively. Besides the ALMAGAL sources, also other classical high-mass hot cores such as Sgr B2(N2) (Belloche et al. 2016), NGC 6334I (Bøgelund et al. 2018), Orion KL (Neill et al. 2013), and CygX-N30 (van der Walt et al. 2021) are included in Figs. 3–4. Only sources where N_{CH₃OH} is derived from the ¹³C or ¹⁸O

isotopologues are included in Fig. 3 to ensure that $N_{\rm CH_3OH}$ is not underestimated. The (D/H)_{CH₃OH} ratios lie mostly in the $10^{-2}-10^{-4}$ range. Interestingly, all the ALMAGAL sources and Orion KL show higher (D/H)_{CH₃OH} ratios ($10^{-2}-10^{-3}$) than Sgr B2(N2) and NGC 6334I ($10^{-3}-10^{-4}$). No clear correlation between the detection of CH₂OH or the derived (D/H)_{CH₃OH} and protostellar parameters such as $L_{\rm bol}$ and envelope mass is present among the high-mass sources. Excluding upper limits, the average (D/H)_{CH₃OH} ratio is $1.1 \pm 0.7 \times 10^{-3}$. This is almost two

orders of magnitude higher than the D/H ratio in the local ISM of $\sim 2 \times 10^{-5}$ (Linsky et al. 2006; Prodanović et al. 2010), suggesting effective deuteration in the cold high-mass prestellar phases. However, both the range of observed (D/H)_{CH₃OH} values and the average is more than one order of magnitude lower than what is generally observed toward low-mass sources (\sim few \times 10⁻², e.g., Bianchi et al. 2017a,b, 2020; Jacobsen et al. 2019; van Gelder et al. 2020, see Sect. 4.1 for further discussion).

Interestingly, the (D/H)_{CH2DOH} ratio (Eq. (3)) is significantly higher than the (D/H)_{CH3OH} ratio, see Fig. 4. For the high-mass sources, only ALMAGAL datapoints are shown since no other interferometric studies of CHD2OH in high-mass protostellar systems are available. The derived (D/H)_{CH2DOH} ratios lie mostly in the 0.1–1 range, with an average of $2.0 \pm 0.8 \times 10^{-1}$, which is more than two orders of magnitude higher than the (D/H)_{CH3OH} ratio. Furthermore, this indicates that about 1/5 of the single deuterated methanol molecules gets successively deuterated further toward CH2DOH in high-mass protostellar systems. This is in good agreement with low-mass protostellar systems where about 1/4 of the CH2DOH is successively deuterated toward CHD2OH (Drozdovskaya et al. 2022).

4. Discussion

4.1. Methanol deuteration from low to high mass

In this work, a (limit on the) (D/H)_{CH3OH} ratio could be derived for 38 of the 99 studied high-mass sources. Since large samples of both low-mass and high-mass protostellar systems with methanol D/H values are now available, a more significant comparison over the mass regime can be made. In Fig. 3, also (D/H)_{CH₃OH} ratios derived for both low-mass prestellar cores and high-mass starless cores are included. It is evident that the (D/H)_{CH₃OH} ratio is lower in high-mass hot cores (10⁻⁴–10⁻²) than in their low-mass counterpart $(10^{-2}-10^{-1})$. Intermediatemass protostars show values in between $(10^{-3}-10^{-2})$, but this subsample only consists of three sources (NGC 7192 FIR2, Cep E-A, and Serpens SMM1-a, Fuente et al. 2014; Ospina-Zamudio et al. 2018; Ligterink et al. 2022, the D/H ratio of NGC 7192 FIR2 is taken from the beam averaged values). However, interestingly the (D/H)_{CH2DOH} ratio seems very similar between low-mass protostars and high-mass protostars (0.1–1, see Fig. 4). Among the low-mass sources, IRAS 2A and IRAS 4A show somewhat elevated (D/H)_{CH2DOH}, but these were derived using older spectroscopic data of CHD₂OH (Taquet et al. 2019).

In Fig. 5, the mean (D/H)_{CH₃OH} ratio is presented for low-mass, intermediate-mass, and high-mass protostellar systems. The average (D/H)_{CH₃OH} ratio for high-mass hot cores (1.1 \pm 0.7 \times 10⁻³) lies more than one order of magnitude lower than the average ratio for low-mass hot corinos (2.2 \pm 1.2 \times 10⁻²), with the average (D/H)_{CH₃OH} for intermediate-mass protostars in between (8.9 \pm 7.4 \times 10⁻³). A similar trend is seen for the high-mass and low-mass prestellar phases where the average (D/H)_{CH₃OH} ratios are 5.9 \pm 5.1 \times 10⁻³ and 3.4 \pm 1.9 \times 10⁻², respectively. The lower (D/H)_{CH₃OH} ratio in both high-mass hot cores and high-mass prestellar phases compared to their lower-mass counterparts suggests a lower deuteration efficiency already in the high-mass prestellar phases (see Sect. 4.3).

The methanol D/H ratios derived in low-mass prestellar cores agree well with those derived for low-mass protostars (see Fig. 3). Since CH₃OH is formed through the hydrogenation of CO ice (e.g., Watanabe & Kouchi 2002; Fuchs et al. 2009; Simons et al. 2020; Santos et al. 2022), this is a strong

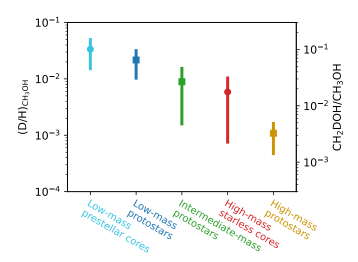


Fig. 5. Average $(D/H)_{CH_3OH}$ ratio derived from N_{CH_2DOH}/N_{CH_3OH} for low-mass, intermediate-mass, and high-mass protostellar systems and low-mass prestellar cores and high-mass starless cores. The prestellar phases are indicated with circles and the protostellar phases with squares.

indication for inheritance of methanol and other COMs ices between low-mass prestellar phases and protostars. On the other hand, the average (D/H)_{CH₃OH} ratio for high-mass starless cores $(5.9 \pm 5.1 \times 10^{-3})$ seems to be about a factor of five higher than that for high-mass protostars $(1.1 \pm 0.7 \times 10^{-3})$. However, the average (D/H)_{CH₃OH} of the high-mass starless cores has a large errorbar since it is based on three detections of which one (G034-G2(MM2)) has a high (D/H)_{CH3OH} ratio of $\sim 10^{-2}$ (Fontani et al. 2015). The other two detections (AFGL 5142-EC and 0548-mm3; Fontani et al. 2015) and all the upper limits show $D/H \lesssim 2 \times 10^{-3}$ which agree well with most of the ALMA-GAL sources as well as with Orion KL (Neill et al. 2013). Only Sgr B2(N2) and most of the cores in NGC 6334I show slightly lower D/H ratios at the 10^{-4} level. This therefore also suggests inheritance of methanol ice from the high-mass prestellar phase to the protostellar phase.

4.2. Singly vs. doubly deuterated methanol

As evident from Figs. 3–4, the methanol D/H ratio derived for CH₂DOH is significantly higher than that derived for CH₃OH. The average (D/H)_{CH₂DOH} ratio is about two orders of magnitude higher (2.0 \pm 0.8 \times 10⁻¹) than (D/H)_{CH₃OH}. In contrast to (D/H)_{CH₃OH}, this is in good agreement with the average of 3.0 \pm 2.0 \times 10⁻¹ for low-mass protostars, suggesting that successive deuteration happens almost equally effective in both low-mass and high-mass systems.

Having higher D/H ratios for the doubly deuterated isotopologue compared with singly deuterated isotopologue is not unique to methanol. For water, the D_2O/HDO ratios are on the order of 10^{-2} (e.g., Coutens et al. 2014; Jensen et al. 2021), which is about an order of magnitude higher than typical HDO/H₂O ratios ($\leq 10^{-3}$; Persson et al. 2014; Jensen et al. 2019; van't Hoff et al. 2022). This difference was attributed to be the result of layered ice chemistry (Dartois et al. 2003; Furuya et al. 2016), where the bulk of the water ice is formed in the warmer translucent cloud phase with a low D/H ratio whereas the surface layers formed in the cold prestellar phases show higher D/H ratios. However, methanol is thought to only start forming in the cold prestellar phases where CO is frozen out

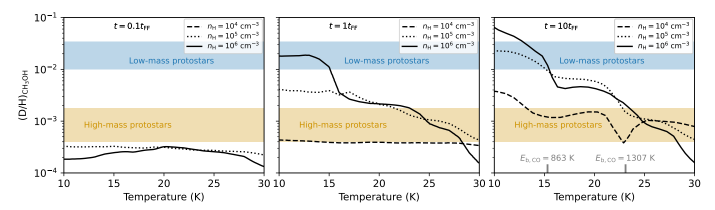


Fig. 6. (D/H)_{CH₃OH} ratio in the ices as function of the gas and dust temperature for $n_{\rm H} = 10^4$ (dashed), 10^5 (dotted), and 10^6 cm⁻³ (solid) as modeled by Bøgelund et al. (2018) and Taquet et al. (2019) using the GRAINOBLE model (Taquet et al. 2012, 2013, 2014). The model results are shown for 0.1 (*left*), 1 (*middle*), and 10 (*right*) times free-fall timescales $t_{\rm FF}$. The observed average (D/H)_{CH₃OH} ratios are indicated in blue and orange for the low-mass and high-mass protostars, respectively. In the *right panel*, the range of possible desorption temperature of CO ice is indicated with the gray bars for binding energies between 863 K and 1307 K for CO ice deposited on non-porous amorphous solid water (Noble et al. 2012).

(e.g., Watanabe & Kouchi 2002; Fuchs et al. 2009) with little to no formation in the warmer translucent phases. Indeed, also for a direct precursor of CH₃OH on the surface of dust grains, H₂CO, the D₂CO/HDCO ratio in IRAS 16293-2422 points toward a high D/H ratio of ~25% compared to a much lower D/H ratio derived from HDCO/H₂CO (~3%; Persson et al. 2018). Small variations in temperature in the 10–20 K range can change the D/H ratio of ice mantle species such as methanol (see Sect. 4.3), but this should affect both CH₂DOH and CHD₂OH in a similar way and should therefore not lead to the observed difference.

One possible explanation could be the optical depth of CH₂DOH. In the low-mass source L1551 IRS5, the emission of CH₂DOH (as well as ¹³CH₃OH) was suggested to be optically thick (Bianchi et al. 2020). However, since the (D/H)_{CH₂OH} ratios derived from CH2DOH clearly show lower values in high-mass protostellar systems compared to their lower-mass counterpart (see Sect. 4.1), this does not seem like a viable solution. Very recently, spectroscopic data for ¹³CH₂DOH has become available (Ohno et al. 2022), but these do not yet include a calculation of the partition function and line properties such as A_{ij} . When assuming that the source size is equal to the beam size, the line optical depth of the most constraining transition, $(17_{1,16} e_0 17_{0.17} e_0$, $E_{\rm up} = 336 \, {\rm K}$) is $\tau < 10^{-2}$. Only for source sizes smaller than <0.5" does CH₂DOH become marginally optically thick $(\tau > 0.1)$ for the most line rich sources. Also, the $(D/H)_{CH_2OH}$ ratios where $N_{\text{CH}_3\text{OH}}$ was derived from the possibly optically thick ¹³C isotopologue are on average less than a factor ~3 higher than the $(D/H)_{CH_3OH}$ ratios where N_{CH_3OH} could be derived using the optically thin ¹⁸O isotopologue (see Fig. E.1).

A more realistic explanation is that successive deuteration of molecules is more effective than the first deuteration. This explanation is supported by several laboratory studies performed at low temperatures of 10–20 K (e.g., Nagaoka et al. 2005, 2007; Hidaka et al. 2009). Drozdovskaya et al. (2022) showed that their observed (D/H)_{CH3OH} ratio as derived from CH2DOH for the low-mass binary IRAS 16293-2422 could be well explained by these experiments whereas CHD2OH and CD3OH were overproduced by the experiments. The latter could be the result of the high atomic D/H flux of 0.1 used in the laboratory studies in contrast to the ISM value of $\sim 10^{-5}$, although the atomic D/H ratio is enhanced in cold dense prestellar cores. Assuming that the (D/H)_{CH3OH} ratio is a direct representative of the gaseous atomic D/H ratio available in the prestellar phases (i.e., that H/D

addition reactions are equally effective), a D/H flux of $\sim 10^{-3}$ may be more realistic for high-mass cold dense cores.

4.3. Linking the methanol D/H to the physical conditions during formation

Given the sensitivity of the methanol deuteration process to both temperature and density (i.e., CO freeze-out), the measured methanol D/H ratios are linked to these physical properties during the prestellar phases. To quantify this for the protostellar systems studied in this work, the observed D/H ratios are compared to the astrochemical gas-grain models presented by Bøgelund et al. (2018) and Taquet et al. (2019). These works used the GRAINOBLE model (Taquet et al. 2012, 2013, 2014) to test the effect of the dust and gas temperature T (assumed to be equal) and hydrogen density $n_{\rm H} = n({\rm H}) + 2n({\rm H_2})$ on the resulting (D/H)_{CH₃OH} ratio in the ices during the prestellar phases. In this work, we compare our results to their results and therefore only a brief description of the model is presented.

In GRAINOBLE, the gas-ice chemistry is computed in three phases: the bulk ice, the ice surface layers, and in the gas phase, following the approach initially presented by Hasegawa & Herbst (1993). The model includes both adsorption and desorption reactions and computes the rate equations in each phase. The chemical network used for the gas-phase chemistry is described in Taquet et al. (2014) and includes both ion-neutral chemistry and all molecules relevant for the chemistry of methanol (e.g., CO, HCO, H₂CO). Moreover, the model computes the deuteration of ice species based on laboratory experiments and includes both the hydrogenation (with both H and D atoms) reactions leading to methanol as well as hydrogen/deuterium abstraction reactions in low temperature (~10–15 K) conditions (Hidaka et al. 2009).

The effect of T and $n_{\rm H}$ on the resulting methanol D/H ratio is presented in Fig. 6 as computed by Bøgelund et al. (2018) and Taquet et al. (2019). For a constant temperature and density, the chemistry was evolved over a timescale indicated on the top of each panel, where the free-fall timescale $t_{\rm FF}$ is 4.4×10^5 , 1.4×10^5 , and 4.4×10^4 yr for $n_{\rm H} = 10^4$, 10^5 , and 10^6 cm⁻³, respectively. For the longest timescales ($t = 10t_{\rm FF}$), it is evident that for all densities the methanol D/H ratio drops with increasing temperature. The strongest decrease is seen for 10^6 cm⁻³, where the methanol D/H ratio decreases from $\sim 6 \times 10^{-2}$ for

T=10 K to as low as $\sim 10^{-4}$ when T=30 K. A similar trend is visible for 10^5 cm^{-3} where the D/H ratio decreases from $\sim 2 \times 10^{-2}$ for T=10 K to $\sim 4 \times 10^{-4}$ at $T\sim 30 \text{ K}$. The decreasing D/H ratios for both these densities is the direct consequence of the decrease of atomic deuterium enhancement in Eq. (1) with increasing temperature. For $n_{\rm H}=10^4 \text{ cm}^{-3}$, the methanol D/H ratio also decreases with temperature, but only by a factor $\sim 4 \text{ between } 10\text{--}30 \text{ K}$.

Another interesting trend is that for decreasing timescales (i.e., moving from right to left in Fig. 6), the D/H ratio at a given temperature also drops for all densities. The strongest drops are seen for the higher density cases at low temperatures (<15 K) where the D/H ratio drops two orders of magnitude from $t = 10t_{\rm FF}$ toward $t = 0.1t_{\rm FF}$. This is the direct result of having less time where CO is frozen out and hence less time to deuterate ice species such as methanol. For higher temperatures (T > 20 K), this effect is less evident since significantly less CO freezes out, although a slightly higher binding energy of CO (up to ~1300 K; Noble et al. 2012) could result in CO frozen out till higher temperatures of ~25 K and hence a higher deuteration efficiency also above 20 K. However, even when CO does not freeze out, CO molecules can still land on the grain for a short period and react with H or D atoms toward HCO, H2CO, and eventually (deuterated) methanol. This effect is most efficient for higher densities of 10⁵–10⁶ cm⁻³ and reduces when timescales smaller than the free-fall timescale are considered. For the lowest density of 10⁴ cm⁻³, this effect is most evident since the CO freeze-out timescale is the highest and therefore the methanol deuteration is hampered the most.

Overplotted in Fig. 6 are the observed methanol (D/H)_{CH₃OH} ratios for both low-mass and high-mass protostars. It is evident that the observed (D/H)_{CH₃OH} ratios suggest a different temperature during methanol formation or different prestellar phase lifetimes for low-mass and high-mass protostars. For high-mass protostars, a temperature of >20 K is needed when the density is larger than 10^5 cm⁻³ and the timescale of the high-mass prestellar phase is > $t_{\rm FF}$. A lower temperature of >13 K at low densities of 10^4 cm⁻³ can also explain the observed methanol D/H ratio toward high-mass protostars, but such low densities in the dense high-mass starless phase are unlikely. Alternatively, the temperature in the high-mass prestellar phases can be in the 15 < T < 20 K range with a lifetime of $\lesssim t_{\rm FF}$. For prestellar lifetimes much smaller than the free-fall timescale, any temperature can explain the observed methanol deuteration toward high-mass protostars.

Contrary to the high-mass protostars, the observed $(D/H)_{CH_3OH}$ ratio for low-mass protostars suggests both a temperature of <15 K and a prestellar phase duration longer than $\geq t_{FF}$. Furthermore, the observed methanol D/H ratio for low-mass protostars cannot be explained by a low density of 10^4 cm⁻³ at any modeled timescale.

It is important to note that the methanol D/H ratio is observed in the gas phase with ALMA whereas the GRAINOBLE models predict the ice abundances in the prestellar phases. Several processes can affect the D/H ratio as the ices warm up while infalling toward the protostar (e.g., Ratajczak et al. 2009; Faure et al. 2015). However, one of the likely dominant processes, CH₂DOH formation through H–D substitution in methanol ice (Nagaoka et al. 2005) is included in the model but this does not dominate over hydrogenation of CO.

These results thus suggest that the high-mass prestellar phases are generally either warm ($T \gtrsim 20$ K) or short ($t \lesssim t_{\rm FF}$) while the low-mass prestellar phases are colder (T < 15 K) and long ($t \geq t_{\rm FF}$). The observed methanol D/H ratios toward

high-mass starless cores and low-mass prestellar cores also fit this picture, see Fig. E.2. On the other hand, the spread in observed abundance ratios of nitrogen-bearing COMs suggests that the scatter in timescales of high-mass prestellar phases is rather small and similar to that of low-mass prestellar phases (Nazari et al. 2022), implying that warmer high-mass pre-stellar phases are a more likely explanation. One possible explanation for the discrepancy between the low-mass and high-mass methanol D/H ratios could be that high-mass stars generally form in clusters with other nearby high-mass stars that heat the surrounding cloud which can affect the D/H ratios molecules forming in the ices (e.g., such as seen for water toward lowmass protostars; Jensen et al. 2019). The majority of the sources studied in this work are located in a clustered environments but these do not show significantly lower D/H ratios than high-mass sources that are single sources at our angular resolution. More modeling work similar to those performed by Bøgelund et al. (2018) and Taquet et al. (2019) including CHD₂OH is needed to further test these hypotheses.

5. Conclusion

In this work, (limits on) the D/H ratios of CH_3OH and CH_2DOH are determined for 38 and 26 sources, respectively, out of the 99 studied sources using ALMA observations of CH_2DOH , CHD_2OH , CH_3OH , CH_3OH , and $CH_3^{18}OH$. The derived $(D/H)_{CH_3OH}$ and $(D/H)_{CH_2DOH}$ ratios are compared to each other as well as to other high-mass protostars, low-mass protostars, and both low-mass and high-mass prestellar phases. Furthermore, comparison with the gas-grain chemical code GRAINOBLE links the observed D/H ratios to the temperature during methanol formation and the lifetime of the prestellar phases. The main conclusions of this work are as follows:

- − The (D/H)_{CH₃OH} ratios of the high-mass protostars studied in this work lie in the 10^{-3} − 10^{-2} range. Combining our sample with other high-mass protostars studied with ALMA, an average (D/H)_{CH₃OH} ratio of $1.1 \pm 0.7 \times 10^{-3}$ is derived. This is in good agreement with the (D/H)_{CH₃OH} ratio derived for high-mass starless cores ($5.9 \pm 5.1 \times 10^{-3}$), but is more than an order of magnitude lower than the average (D/H)_{CH₃OH} ratio for low-mass protostars ($2.2 \pm 1.2 \times 10^{-2}$) and low-mass prestellar cores ($3.4 \pm 1.9 \times 10^{-2}$).
- For $(D/H)_{CH_2DOH}$, significantly higher values than $(D/H)_{CH_3OH}$ are found ranging from 0.1–1 with an average of $2.0 \pm 0.8 \times 10^{-1}$. The latter is good agreement with results on low-mass protostars and suggests that about 1/5 singly deuterated methanol molecules gets successively deuterated further independent of the mass of the system.
- Based on a comparison with GRAINOBLE models in the literature, the lower (D/H)_{CH₃OH} ratios toward high-mass protostars suggest either a temperature of \gtrsim 20 K in the high-mass prestellar phases or a short lifetime ($\lesssim t_{FF}$) of the high-mass prestellar phases. This is in strong contrast with the low-mass sources for which the higher (D/H)_{CH₃OH} ratio can only be achieved when the low-mass prestellar phases are both cold (<15 K) and long lived ($\ge t_{FF}$).

This work demonstrates that the deuteration of the CH₃-group of methanol as measured toward protostellar systems could be used to probe the physical conditions (e.g., temperature) of the prestellar phases. The discrepancy in (D/H)_{CH₃OH} between low-mass and high-mass sources indicates that the physical conditions are already different before the onset of star formation. Additional observations of multiple deuterated methanol isotopologues (e.g., CHD₂OH, CD₃OH) as well as CH₃OD

will shed further light on the efficiency of methanol deuteration between low-mass and high-mass systems. In combination with additional modeling studies (such as those performed by Bøgelund et al. 2018; Taquet et al. 2019; Kulterer et al. 2022), this can provide further insight on the relevant deuterium chemistry and how the D/H ratio varies across the protostellar mass range.

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Appendix A: Transitions of CH₃OH and isotopologues

Table A.1. Transitions of CH₃OH and isotopologues with $A_{ij} > 10^{-6}$ and $E_{\rm up} < 1000$ K covered in the ALMAGAL (2019.1.00195.L) program.

Species		nsit		Frequency	A_{ij}	$E_{ m up}$
Species	(J K L M	-	JKLM)	(GHz)	(s^{-1})	(K)
CH ₃ OH	6153	_	7253	217.2992	4.3×10^{-5}	373.9
- 3-	15 6 9 3	_	16 5 11 3	217.6427	1.9×10^{-5}	745.6
	15 6 10 3	_	16 5 12 3	217.6427	1.9×10^{-5}	745.6
	20 1 19 1	_	20 0 20 1	217.8865	3.4×10^{-5}	508.4
	4231	_	3 1 2 1	218.4401	4.7×10^{-5}	45.5
	25 3 23 1	_	24 4 20 1	219.9837	2.0×10^{-5}	802.2
	23 5 18 1	_	22 6 17 1	219.9937	1.7×10^{-5}	775.9
	8081	_	7161	220.0786	2.5×10^{-5}	96.6
	10 5 6 2	_	11 4 8 2	220.4013	1.1×10^{-5}	251.6
¹³ CH ₃ OH	14 1 13 -0	_	13 2 12 -0	217.0446	2.4×10^{-5}	254.3
011,011	10 2 8 +0	_	937+0	217.3995	1.5×10^{-5}	162.4
	17 7 11 +0	_	18 6 12 +0	220.3218	1.3×10^{-5}	592.3
	17 7 10 -0	_	18 6 13 -0	220.3218	1.3×10^{-5}	592.3
CH ₃ ¹⁸ OH	14 1 14 1	-	13 2 12 1	217.1729	1.7×10^{-5}	238.9
C11 ₃ C11	18 6 13 4	_	17 7 11 4	217.9223	1.5×10^{-5}	874.1
	17 5 13 4	_	18 6 13 4	218.5521	3.2×10^{-5}	884.6
	4222	_	3 1 2 2	219.4078	4.6×10^{-5}	44.6
	8715	_	7615	219.8433	2.8×10^{-5}	663.2
	18 3 16 5	_	19 4 16 5	219.9572	5.1×10^{-5}	795.8
	8181	_	7071	220.1951	3.6×10^{-5}	85.7
CH ₂ DOH	26 4 22 0	_	26 3 24 2	217.2664	2.0×10^{-5}	817.1
CH ₂ DOH	26 1 25 2	_	26 1 26 2	217.2004	1.1×10^{-5}	777.9
	17 4 13 2	_	16 5 11 1	217.3436	5.2×10^{-6}	409.7
	17 4 13 2	_	16 5 12 1	217.3593	5.2×10^{-6} 5.3×10^{-6}	409.7
	23 6 18 1	_	22 7 15 0	217.3818	6.4×10^{-6}	742.6
	23 6 17 1	_	22 7 16 0	217.3825	6.4×10^{-6}	742.6
	18 1 17 2	_	18 2 17 0	217.4479	1.8×10^{-5}	391.5
	25 1 25 2	_	25 0 25 1	217.6429	4.7×10^{-5}	712.4
	12 7 6 0	_	13 6 7 1	217.6446	2.5×10^{-6}	357.2
	12 7 5 0	_	13 6 8 1	217.6446	2.5×10^{-6} 2.5×10^{-6}	357.2
	18 0 18 0	_	17 1 16 2	218.1095	8.9×10^{-6}	363.2
	5 2 4 1	_	5 1 5 1	218.3164	9.1×10^{-6}	58.7
	24 3 21 1	_	24 2 23 2	218.5348	3.9×10^{-5}	687.7
	20 5 16 1	_	19 6 13 0	219.2043	1.5×10^{-5}	557.6
	20 5 15 1	_	19 6 14 0	219.2061	1.5×10^{-5} 1.5×10^{-5}	557.6
	5 1 5 1	_	4141	219.5515	7.0×10^{-6}	48.2
	5151	_	4140	220.0718	3.3×10^{-5}	35.8
	29 4 26 0	_	29 3 26 2	220.3492	2.5×10^{-5}	997.1
	17 1 16 0	_	17 0 17 0	220.5492	2.3×10^{-5} 3.8×10^{-5}	335.9
	21 1 20 1	_	21 1 21 1	220.5326	2.0×10^{-6}	515.1
	21 2 19 2	_	21 1 21 1 21 1 21 1 20 2	220.0230	3.9×10^{-5}	531.1
CHD ₂ OH	6221	_	5122	217.0702	$\frac{3.9 \times 10^{-6}}{1.8 \times 10^{-6}}$	61.9
СПРЗОП	13 2 1 2	-	12 3 1 2	217.0702	8.8×10^{-6}	213.0
	16 5 1 2	_	12 3 1 2	217.1181	2.4×10^{-6}	366.8
	7012	_	6112	217.2031 217.4912	2.4×10^{-5} 2.9×10^{-5}	74.3
	7421	_	8320	217.4912	2.9×10^{-6} 4.4×10^{-6}	74.3 111.1
	7421	-	8320		4.4×10^{-6} 4.4×10^{-6}	111.1
		-		217.5430	4.4×10^{-6} 9.0×10^{-6}	854.9
	24 9 1 1		25 8 1 0	217.8034	9.0×10^{-6} 9.0×10^{-6}	
	24 9 2 1	-	25 8 2 0	217.8034		854.9
	2211	-	3110	218.0092	5.5×10^{-6}	25.8
	7210	-	7120	218.1279	1.2×10^{-5}	68.8
	16 3 2 2	-	15 4 2 2	218.2323	6.9×10^{-6}	318.2
	20 2 2 2	-	20 2 1 1	218.4156	9.0×10^{-6}	449.9

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Table A.1. continued.

Species	Tra	nsit	ion	Frequency	A_{ij}	$E_{ m up}$
	(J K L M)	-	JKLM)	(GHz)	(s^{-1})	(K)
	11 2 1 2	-	11 1 2 2	218.4824	2.3×10^{-5}	163.0
	12 2 1 2	-	12 1 2 2	219.2181	4.9×10^{-6}	187.0
	23 10 2 2	-	24 9 2 2	219.3323	5.1×10^{-6}	872.6
	23 10 1 2	-	24 9 1 2	219.3323	5.1×10^{-6}	872.6
	16 3 1 0	-	15 4 1 0	219.3377	6.5×10^{-6}	300.9
	22 5 2 0	-	22 4 1 2	219.6514	2.6×10^{-6}	583.5
	22 5 1 0	-	22 4 2 2	219.7983	2.5×10^{-6}	583.4
	13 2 1 0	-	12 3 1 0	220.2430	4.7×10^{-6}	195.3
	11 6 1 1	-	12 5 1 1	220.5567	1.1×10^{-6}	249.5
	11 6 2 1	-	12 5 2 1	220.5569	1.1×10^{-6}	249.5

Notes. The typical beam size is $\theta_{beam} \sim 1''$ and the typical rms is rms_{line} ~ 0.2 K.

Appendix B: Observational details

Table B.1. Column densities of ¹³CH₃OH, CH₃⁸OH, CH₃OH, CH₂DOH, and CHD₂OH and derived methanol D/H ratios.

" K 18:26:48:92 -12:26:24:51 1.23 150 18:26:47:96 -12:26:20:73 1.23 150 18:26:47:96 -12:26:25:98 1.23 150 18:25:00:82 -13:15:34:46 1.22 150 18:25:01:01 -13:15:38.57 1.22 150 18:25:01:05 -12:42:21:16 1.22 150 18:25:10:06 -12:42:21:16 1.23 150 18:25:10:06 -12:42:21:16 1.23 150 18:25:10:06 -12:42:21:14 1.23 150 18:25:10:06 -12:42:21:14 1.23 150 18:25:10:06 -12:42:24.74 1.23 150 18:25:10:06 -12:42:21:14 1.23 150 18:25:10:06 -12:42:24.74 1.23 150 18:25:10:06 -12:42:24.74 1.23 150 18:25:10:05 -12:42:24.74 1.23 150 18:34:23:39 -09:00:38:44 1.25 150 18:34:25:55 -	155 155 155 155 155 155 155 155		cm ⁻² <2.1(15) <1.5(15) <1.7(15) <2.5(15) <2.5(15) <3.1(15) <3.6(15) <1.7(15) <1.7(15) <1.7(15) <1.7(15) <1.7(15) <2.4(15) <3.0(15) <3.0(15) <3.0(15) <3.0(15) <3.0(15) <3.0(15) <4.2(15) <3.5(15) <4.2(15) <4.2(15) <4.2(15) <4.2(15)	cm ⁻² <2.4(15) <4.0(15) <4.0(15) <4.0(15) <4.7(15) <2.2(15) <4.0(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.4(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(15) <5.2(
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08:32:09.06	<2.9(15) <1.6	<1.6(15) $0.1-2.1(17)$	<5.6(15)	<4.3(15)	I	I
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	_	7	<4.0(15)	<3.8(15)	<6.7(-3)	1
644284B 10:31:29.63 -58:02:18.82 0.86 100 8.2±1.	$8.2\pm1.8(15)$ $1.5\pm$	$(.5\pm0.3(15) 7.8\pm4.3(17)$	<3.0(15)	<3.9(15)	<2.9(-3)	1
12:35:35.05 -63:02:31.19 0.99 200	(9)	(5)	$1.1\pm0.3(16)$	<3.0(15)	$2.7\pm1.7(-3)$	<3.9(-1)
150		<1.2(15) <1.8(15)	<5.9(15)	<7.9(15)	I	I
13:11:14.14 -62:45:06.80 1.29 150		•	<4.5(15)	<2.2(15)	1	1
13:12:36.17 -62:33:34.43 0.87 150	15)	$3.0\pm1.5(15)$ $1.3\pm0.8(18)$	<6.3(15)	<4.9(15)	<3.7(-3)	1
	<2.7(15) <1.8	<1.8(15) 0.2–16.4(16)	<4.8(15)	<2.9(15)	I	1
13:14:22.99 -62:45:54.35 0.87 150	·		<6.6(15)	<2.9(15)	I	I
13:14:23.07 -62:45:47.54 0.87 150	·		<4.9(15)	<9.0(15)	I	1
13:14:26.92 -62:44:29.72 0.88 150	•	$(15) \qquad 0.1-1.9(17)$	<6.2(15)	<2.6(15)	ı	ı
706785B 13:14:26.55 -62:44:31.80 0.88 150 <3.3(1	<3.3(15) <2.9	<2.9(15) 0.0–2.0(17)	<6.0(15)	<3.6(15)	1	1

Table B.1. continued.

Source	RA (J2000)	Dec (J2000)	$\theta_{ m beam}$	$T_{\rm ex}$	$N_{^{13}\mathrm{CH_{3}OH}}$	$N_{ m CH^{18}OH}$	<i>N</i> сн ₃ он	N _{СН2} DОН	$N_{\text{CHD}_2\text{OH}}$	$(D/H)_{CH_3OH}$	$(D/H)_{CH_2DOH}^2$
			:	×	cm^{-2}	cm^{-2}	cm^{-2}	cm^{-2}	cm^{-2}		
706785C	13:14:26.38	-62:44:30.24	0.88	150	<2.7(15)	<1.3(15)	<1.7(15)	<5.9(15)	<9.0(15)	1	1
706785D	13:14:25.64	-62:44:30.36	0.88	150	<4.4(15)	<3.2(15)	0.1 - 2.7(17)	<6.1(15)	<3.7(15)	1	1
707948	13:16:43.19	-62:58:32.83	0.88	150	$1.8\pm0.4(17)$	$1.5\pm0.3(16)$	$1.1\pm0.6(19)$	$2.0\pm1.0(17)$	$6.5\pm4.8(16)$	$6.0\pm 4.3(-3)$	$3.2\pm 2.9(-1)$
717461A	13:43:01.68	-62:08:51.42	1.29	150	$7.6\pm1.9(15)$	$2.6\pm0.8(15)$	$1.1\pm0.4(18)$	<3.7(15)	<2.5(15)	<2.0(-3)	ı
717461B	13:43:01.74	-62:08:55.34	1.29	150	<2.2(15)	<1.9(15)	<2.8(15)	<3.8(15)	<3.6(15)	I	I
721992	13:51:58.27	-61:15:41.04	0.85	150	<4.4(15)	<1.9(15)	0.5-2.5(17)	<7.3(15)	<5.5(15)	1	I
724566	13:59:30.92	-61:48:38.27	0.83	150	$2.9\pm0.6(16)$	$1.0\pm0.2(16)$	$4.0\pm 2.3(18)$	$2.0\pm1.0(16)$	<9.6(15)	$1.7\pm1.3(-3)$	<9.6(-1)
732038	14:13:15.05	-61:16:53.19	0.82	150	<7.6(15)	<4.8(15)	1.2-4.2(17)	<6.8(15)	<7.5(15)	ı	ı
7447 <i>5</i> 7A	14:45:26.35	-59:49:15.55	1.30	150	$1.4\pm0.4(16)$	$4.1\pm0.9(15)$	$1.7\pm1.0(18)$	$8.0\pm4.0(15)$	<2.5(15)	$1.6\pm1.2(-3)$	<6.3(-1)
744757B	14:45:26.16	-59:49:19.87	1.30	150	<1.9(15)	<4.1(15)	1.3–11.3(16)	<2.9(15)	<2.3(15)	1	1
759150A	15:10:43.13	-57:44:49.63	1.29	150	<1.9(15)	<1.2(15)	0.3 - 10.1(16)	<5.0(15)	<2.3(15)	I	I
759150B	15:10:43.52	-57:44:44.82	1.29	150	<2.4(15)	<9.6(14)	<1.5(15)	<5.0(15)	<2.8(15)	ı	ı
759150C	15:10:44.48	-57:44:47.33	1.29	150	<2.2(15)	<9.6(14)	<8.6(14)	<5.0(15)	<2.3(15)	ı	ı
759150D	15:10:42.71	-57:44:52.85	1.29	150	<5.0(15)	<9.7(14)	<1.2(15)	<4.9(15)	<2.3(15)	I	I
759150E	15:10:44.10	-57:44:52.03	1.29	150	<2.1(15)	<9.6(14)	<9.9(14)	<4.8(15)	<2.3(15)	1	I
767784	15:29:19.31	-56:31:22.02	1.29	100	$3.9\pm0.5(16)$	$7.2\pm1.6(15)$	$2.5\pm1.4(18)$	$1.0\pm0.4(16)$	<3.3(15)	$1.4\pm1.0(-3)$	<5.3(-1)
800287	16:11:26.57	-51:41:57.14	08.0	100	$1.5\pm0.3(16)$	$6.2\pm1.4(15)$	$1.8\pm1.1(18)$	$1.0\pm0.5(16)$	<1.3(16)	$1.9\pm1.5(-3)$	<2.7(0)
854214A	16:52:32.74	-43:23:49.60	1.26	150	<2.2(15)	<1.7(15)	0.7 - 9.5(16)	<3.8(15)	<5.0(15)	I	I
854214B	16:52:33.02	-43:23:50.26	1.26	150	<5.0(15)	<7.8(14)	0.1-2.1(17)	<3.8(15)	<2.7(15)	1	I
863312A	17:02:08.36	-41:46:56.89	0.83	150	<8.9(15)	<3.7(15)	<3.8(15)	<2.0(16)	<1.4(16)	I	I
863312B	17:02:09.14	-41:46:45.04	0.83	150	<3.7(15)	<1.9(15)	<2.0(15)	<6.0(15)	<4.2(15)	1	I
865468A C1	17:05:10.90	-41:29:06.99	1.23	100	$1.3\pm0.3(17)$	$3.1\pm0.7(16)$	$1.0\pm0.6(19)$	$1.5\pm0.8(17)$	$4.0\pm 2.0(16)$	$4.9\pm3.8(-3)$	$2.7 \pm 1.9(-1)$
865468A C2	17:05:10.90	-41:29:06.99	1.23	150	$5.1\pm1.1(16)$	<3.0(15)	$2.5\pm1.4(18)$	$6.0\pm3.0(16)$	<2.0(16)	$7.9\pm5.7(-3)$	<6.7(-1)
865468B	17:05:11.22	-41:29:07.65	1.24	150	$1.3\pm0.2(16)$	$2.6\pm0.6(15)$	$8.6\pm5.0(17)$	$6.5\pm2.0(15)$	<5.0(15)	$2.5\pm1.7(-3)$	<1.1(0)
865468C	17:05:11.12	-41:29:03.47	1.24	150	$5.0\pm1.9(15)$	$1.0\pm0.2(15)$	$3.4\pm2.0(17)$	<7.5(15)	<5.8(15)	<1.8(-2)	1
876288	17:11:51.02	-39:09:29.18	0.81	150	$8.9\pm2.9(15)$	$3.1\pm0.7(15)$	$5.7\pm3.9(17)$	<6.0(15)	<5.0(15)	<1.2(-2)	I
881427A	17:20:06.31	-38:57:15.18	1.23	150	$9.3\pm1.8(16)$	$2.6\pm0.7(16)$	$1.1\pm0.4(19)$	$8.3\pm1.6(16)$	$1.9\pm0.7(16)$	$2.5\pm1.1(-3)$	$2.3\pm1.0(-1)$
881427B	17:20:06.46	-38:57:11.44	1.23	300	$1.0\pm0.2(16)$	$8.2\pm1.8(15)$	$3.4\pm1.9(18)$	$1.5\pm0.8(16)$	<1.0(16)	$1.5\pm1.1(-3)$	<1.3(0)
881427C	17:20:06.12	-38:57:15.84	1.23	150	$7.4\pm1.6(16)$	$2.6\pm0.5(16)$	$1.1\pm0.4(19)$	$5.5\pm1.2(16)$	$1.2\pm0.5(16)$	$1.7\pm0.7(-3)$	$2.2\pm0.9(-1)$
G023.3891+00.1851	18:33:14.32	-08:23:57.82	1.24	200	$7.9\pm2.2(15)$	$2.5\pm1.2(15)$	$7.9\pm4.5(17)$	$9.5\pm3.0(15)$	<5.0(15)	$4.0\pm 2.6(-3)$	<7.6(-1)
G023.6566-00.1273	18:34:51.57	-08:18:21.81	1.24	150	$1.4\pm0.4(16)$	<5.0(15)	$7.1\pm2.5(17)$	$2.5\pm1.2(16)$	<5.0(15)	$1.2\pm0.7(-2)$	<4.0(-1)
G025.6498+01.0491	18:34:20.92	-05:59:42.08	1.17	150	$3.0\pm0.6(16)$	$8.5\pm2.7(15)$	$3.4\pm1.4(18)$	$1.5\pm0.8(16)$	$5.0\pm2.5(15)$	$1.5\pm1.0(-3)$	$3.3\pm2.4(-1)$
G030.1981-00.1691	18:47:03.05	-02:30:36.30	09.0	150	$5.1\pm1.1(15)$	<1.8(15)	$2.2\pm1.2(17)$	<1.5(16)	<1.6(16)	<4.7(-2)	1
G233.8306-00.1803	07:30:16.73	-18:35:49.06	0.81	150	<3.7(15)	<3.0(15)	<2.6(15)	<6.5(15)	<4.4(15)	1	1
G305.2017+00.2072A1	13:11:10.45	-62:34:38.60	1.30	150	$1.2\pm0.3(16)$	$2.0\pm0.5(15)$	$8.5\pm4.8(17)$	$5.0\pm 2.5(15)$	<2.3(15)	$2.0\pm1.5(-3)$	<9.1(-1)
G305.2017+00.2072A2	13:11:13.12	-62:34:42.74	1.30	150	<8.8(15)	<5.0(15)	0.1 - 5.3(17)	<8.0(15)	<1.2(16)	I	ı
G310.0135+00.3892	13:51:37.88	-61:39:07.74	1.30	150	<2.3(15)	<2.6(15)	0.3–13.5(16)	<7.9(15)	<2.9(15)	I	ı
G314.3197+00.1125	14:26:26.25	-60:38:31.26	1.30	150	<2.4(15)	<2.2(15)	0.8 - 1.4(17)	<1.0(16)	<6.0(15)	ı	1
G316.6412-00.0867	14:44:18.35	-59:55:11.28	1.29	100^{3}	$2.8\pm0.4(16)$	$8.6\pm2.7(15)$	$3.5\pm1.5(18)$	$2.8\pm0.5(16)$	$4.8\pm3.0(15)$	$2.7\pm1.2(-3)$	$1.7\pm1.1(-1)$
G318.0489+00.0854B	14:53:42.64	-59:08:53.02	1.30	150	$1.7\pm0.3(16)$	$4.9\pm1.9(15)$	$1.9\pm0.9(18)$	$8.0\pm 4.0(15)$	<3.0(15)	$1.4\pm1.0(-3)$	<7.5(-1)
G318.9480-00.1969A1	15:00:55.28	-58:58:52.60	1.29	100^{3}	$8.5\pm1.2(16)$	$2.2\pm0.4(16)$	$9.6\pm 3.2(18)$	$7.0\pm1.1(16)$	$1.4\pm0.7(16)$	$2.4\pm0.9(-3)$	$2.0\pm1.1(-1)$
G318.9480-00.1969A2	15:00:55.23	-58:58:55.88	1.29	150	<2.6(15)	<7.8(14)	0.4 - 16.0(16)	<6.5(15)	<2.3(15)	ı	ı
G323.7399-00.2617B1	15:31:45.64	-56:30:50.16	1.28	150	8.7±2.0(15)	2.0±0.5(15)	7.5±4.3(17)	<2.8(15)	<3.0(15)	<2.9(-3)	1

Table B.1. continued

Source	RA (J2000)	RA (J2000) Dec (J2000)	$\theta_{ m beam}$	$T_{ m ex}$	N ₁₃ СH ₃ OH сm ⁻²	N _{CH} ¹⁸ Oн сm ⁻²	$N_{\rm CH_3OH}$ cm ⁻²	$N_{\mathrm{CH}_2\mathrm{DOH}}$ cm $^{-2}$	$N_{\mathrm{CHD}_2\mathrm{OH}}$	(D/H) _{CH3} OH	$(\mathrm{D/H})_{\mathrm{CH}_2\mathrm{DOH}}^2$
G323.7399-00.2617B2	15:31:45.45	-56:30:49.84	1.28	125	8.2±1.8(16)	$1.5\pm0.3(16)$	5.7±3.3(18)	$2.0\pm1.0(16)$	$4.0\pm2.0(15)$	$1.2\pm0.9(-3)$	2.0±1.4(-1)
G323.7399-00.2617B3	15:31:45.73	-56:30:51.93	1.28	150	<2.5(15)	<6.0(14)	1.4–13.4(16)	<4.7(15)	<2.3(15)	I	I
G323.7399-00.2617B4	15:31:45.94	-56:30:51.34	1.28	150	<2.6(15)	<6.9(14)	1.2 - 14.2(16)	<2.7(15)	<2.8(15)	I	I
G323.7399-00.2617B5	15:31:45.62	-56:30:45.62	1.28	150	<2.6(15)	<5.8(14)	0.1 - 13.9(16)	<5.0(15)	<2.7(15)	I	ı
G323.7399-00.2617B6	15:31:45.84	-56:30:47.68	1.28	150	<3.9(15)	<1.2(15)	0.1-2.1(17)	<2.6(15)	<3.7(15)	ı	ı
G323.7399-00.2617B7	15:31:45.91	-56:30:46.10	1.28	150	<1.7(15)	<1.2(15)	0.1-9.3(16)	<3.9(15)	<1.6(15)	I	ı
G327.1192+00.5103	15:47:32.72	-53:52:38.60	0.81	100	$4.0\pm0.9(16)$	$9.2\pm2.0(15)$	$2.9\pm1.7(18)$	$3.5\pm1.8(16)$	<5.0(15)	$4.0\pm3.1(-3)$	<2.9(-1)
G343.1261-00.0623	16:58:17.22	-42:52:07.54	1.25	100	$8.8\pm2.7(15)$	<7.0(15)	$5.0\pm1.8(17)$	<4.0(16)	$4.0\pm 2.0(15)$	<4.1(-2)	>5.0(-2)
G345.5043+00.3480 C1	17:04:22.89	-40:44:23.06	1.25	125	$5.1\pm1.1(16)$	$1.0\pm0.2(16)$	$4.0\pm 2.3(18)$	$5.0\pm2.5(16)$	$1.0\pm0.5(16)$	$4.2\pm3.2(-3)$	$2.0\pm1.4(-1)$
G345.5043+00.3480 C2	17:04:22.89	-40:44:23.06	1.25	150	$3.1\pm0.7(16)$	<1.0(16)	$1.7\pm0.9(18)$	$3.0\pm1.5(16)$	<1.0(16)	$5.9\pm4.3(-3)$	<6.7(-1)
G348.7342-01.0359B1	17:20:07.08	-38:57:11.22	1.23	150	<3.4(15)	<9.9(14)	1.3–17.7(16)	<1.2(16)	<3.9(15)	I	I
G348.7342-01.0359B2	17:20:07.26	-38:57:09.82	1.23	150	<2.3(15)	<1.7(15)	0.1 - 11.9(16)	<3.6(15)	<4.1(15)	ı	ı
G348.7342-01.0359B3	17:20:07.38	-38:57:10.15	1.23	150	<4.0(15)	<1.5(15)	<1.4(15)	<1.1(16)	<3.8(15)	1	1

(i.e., beam dilution = 1). The column density of CH₃OH is derived from the CH₃OH and ¹³CH₃OH and ¹³CH₃OH isotopologues when these are detected. If neither isotopologue is detected but CH₃OH is, a Notes. The coordinates mark the position from which the spectra were extracted. All column densities are derived for the reported T_{cs} and assuming the source size is equal to the size of the beam range in column densities is presented where the lower limit it the column density derived from CH₃OH and the upper limit is scaled from the upper limit on ¹³CH₃OH. If CH₃OH itself is also not detected, an upper limit is directly derived from its spectrum.

(i) Derived from $N_{\text{CH}_2\text{OH}}$ using Eq. (2). (3) Eq. (3). (3) T_{ex} is set to 150 K for deuterated isotopologues and to 100 K for the other isotopologues.

Appendix C: Methanol D/H ratios of sources in the literature

Table C.1. $(D/H)_{CH_3OH}$ and $(D/H)_{CH_2DOH}$ ratios taken from the literature that are included in Figs. 3 and 4 .

Source	Type ⁽¹⁾	$(D/H)_{CH_3OH}$	$(D/H)_{CH_2DOH}$	Refs
В1-с	LMP	2.8±0.9(-2)	1.3±0.2(-1)	1,2
Serpens S68N	LMP	$1.4\pm0.6(-2)$	$1.2\pm0.5(-1)$	1,2
B1-bS	LMP	<1.8(-2)	_	1,2
HH212	LMP	$3.8\pm2.3(-2)$	_	3
		$8.1\pm3.0(-3)$	_	4
IRAS 16293A	LMP	$2.8 \pm 1.2(-2)$	$2.0\pm0.7(-1)$	5,6
IRAS 16293B	LMP	$2.4 \pm 0.9(-2)$	$2.5\pm0.9(-1)$	6,7
IRAS 2A	LMP	$1.9 \pm 1.0(-2)$	$7.0\pm2.6(-1)$	8
IRAS 4A	LMP	$1.4\pm0.8(-2)$	$5.6 \pm 2.2(-1)$	8
L483	LMP	$7.8\pm3.3(-3)$	_	9
BHR71	LMP	$9.6 \pm 4.1(-3)$	_	10
Ser-emb 1	LMP	<2.0(-1)	_	11
Ser-emb 11W	LMP	<4.3(-2)	_	12
HOPS-108	LMP	$7.0\pm 3.8(-3)$	_	13
G192.12-11.10	LMP	$9.8 \pm 5.8(-3)$	_	14
G205.46-14.56S1-A	LMP	$6.7\pm3.3(-3)$	_	14
G208.68-19.20N1	LMP	$1.3\pm0.6(-2)$	_	14
G210.49-19.79W-A	LMP	$1.4\pm0.6(-2)$	_	14
G211.47–19.27S	LMP	$1.7\pm0.4(-2)$	_	14
V883 Ori	LMP	$4.8 \pm 1.0(-2)$	_	15
Serpens SMM1-a	IMP	$3.6\pm2.1(-3)$	_	16
NGC 7192 FIRS2	IMP	1.9±0.8(-3)	_	17
Cep E-A	IMP	1.2±0.4(-2)	_	18
Sgr B2(N2)	HMP	$4.0\pm1.7(-4)$	_	19,20
NGC6334I MM1 I	HMP	$3.0\pm1.7(-4)$	_	21
NGC6334I MM1 II	HMP	1.8±0.9(-4)	_	21
NGC6334I MM1 III	HMP	$3.2 \pm 1.8(-4)$	_	21
NGC6334I MM1 IV	HMP	1.6±1.0(-4)	_	21
NGC6334I MM1 V	HMP	$2.3\pm1.1(-4)$	_	21
NGC6334I MM2 I	HMP	6.0±3.6(-4)	_	21
NGC6334I MM2 II	HMP	$1.5\pm0.7(-4)$	_	21
NGC6334I MM3 I	HMP		_	21
NGC6334I MM3 II	HMP	$2.4 \pm 1.1(-4)$		21
		$2.6 \pm 1.1(-4)$	_	22
Orion KL ridge	HMP	1.7±0.7(-3)	_	
Orion KL HC	HMP	<1.7(-3)	_	22
CygX-N30	HMP	<1.7(-3)	_	23 24
L1495-B10 6	LMPC LMPC	$1.4 \pm 0.4(-2)$		24 24
L1495-B10 7		$4.8 \pm 1.3(-2)$	_	
L1495-B10 8	LMPC	$5.2\pm1.5(-2)$	_	24
L1495-B10 9	LMPC	$4.6 \pm 1.1(-2)$	_	24
L1495-B10 10	LMPC	$4.2\pm1.3(-2)$	_	24
L1495-B10 11	LMPC	<1.9(-2)	_	24
L1495-B10 12	LMPC	$2.0\pm0.4(-2)$	_	24
L1495-B10 13	LMPC	<3.5(-2)	_	24
L1495-B10 14	LMPC	<1.4(-2)	_	24
L1495-B10 15	LMPC	$4.0\pm1.1(-2)$	_	24
L1495-B10 16	LMPC	$3.0\pm1.0(-2)$	_	24
L1495-B10 17	LMPC	$8.8 \pm 2.7(-2)$	_	24
L183	LMPC	$1.3\pm0.2(-2)$	_	25
L1544	LMPC	$3.0\pm1.3(-2)$	_	26
I00117-MM2	HMSC	<2.4(-3)	_	27
AFGL 5142-EC	HMSC	$9.0 \pm 3.8(-4)$	_	27
05458-mm3	HMSC	$2.3\pm1.0(-3)$	_	27
G034-G2(MM2)	HMSC	$1.4\pm0.6(-2)$	_	27
G034-F2(MM7)	HMSC	<2.5(-3)	_	27
G034-F1(MM8)	HMSC	<1.5(-3)	_	27

Table C.1. continued.

Source	Type ^(I)	(D/H) _{CH₃OH}	(D/H) _{CH₂DOH}	Refs
G034-C1(MM9)	HMSC	<8.7(-4)	_	27
I20293-WC	HMSC	<1.9(-3)	_	27
I22134-G	HMSC	<1.7(-3)	_	27
I22134-B	HMSC	<6.7(-3)	_	27

Notes. The $(D/H)_{CH_3OH}$ and $(D/H)_{CH_2DOH}$ ratios are either directly taken from the reported literature or computed using Eqs. (2) and (3) using the column densities from the reported literature. A 30% uncertainty was assumed in cases where no uncertainty was reported. (1) LMP: low-mass protostar, IMS: intermediate-mass protostar, HMS: high-mass protostar, LMPC: low-mass prestellar core, HMSC: high-mass starless core.

References. 1: van Gelder et al. (2020); 2: Appendix D; 3: Lee et al. (2019a); 4: Bianchi et al. (2017b); 5: Manigand et al. (2020); 6: Drozdovskaya et al. (2022); 7: Jørgensen et al. (2018); 8: Taquet et al. (2019); 9: Jacobsen et al. (2019); 10: Yang et al. (2020); 11: Martín-Doménech et al. (2019); 12: Martín-Doménech et al. (2021); 13: Chahine et al. (2022); 14: Hsu et al. (2022); 15: Lee et al. (2019b); 16: Ligterink et al. (2021); 17: Fuente et al. (2014); 18: Ospina-Zamudio et al. (2018); 19: Belloche et al. (2016); 20: Müller et al. (2016); 21: Bøgelund et al. (2018); 22: Neill et al. (2013); 23: van der Walt et al. (2021); 24: Ambrose et al. (2021); 25: Lattanzi et al. (2020); 26: Bizzocchi et al. (2014); 27: Fontani et al. (2015).

Appendix D: Doubly deuterated methanol in B1-c, Serpens S68N, and B1-bS

Using the database entry of CHD₂OH provided by Drozdovskaya et al. (2022), transitions from CHD₂OH can also be searched for in a few COM-rich low-mass protostars. Here, this is done for B1-c, Serpens S68N (hereafter S68N), and B1-bS from the 2017.1.01174.S ALMA program. The content of oxygen-bearing COMs for these sources was presented by van Gelder et al. (2020), but no public database entry was yet available for CHD₂OH at that time.

Only one strong transition of CHD₂OH ($7_{1,2}$ o_1 – $7_{0,1}$ e_0 , $E_{\rm up}$ = 68 K) is available for these sources which lies on the very edge of the covered frequency range. For B1-c and S68N, this transition is detected at the 3σ level, but given that only half the line is observed this detection is still tentative. Using the same method for deriving the column densities as used by van Gelder et al. (2020) and assuming an excitation temperature of 150 K, we derive column densities of $2.0 \pm 0.6 \times 10^{16}$ cm⁻² for B1-c, $7.2 \pm 2.7 \times 10^{15}$ cm⁻² for S68N, and $< 1.7 \times 10^{15}$ cm⁻² for B1-bS. The FWHM was fixed to the average FWHM of those sources of 3.2 km s⁻¹, and 1.0 km s⁻¹, respectively (van Gelder et al. 2020). The resulting fits are shown in Fig. D.1. Using these derived column densities and those reported for CH₃OH by van Gelder et al. (2020), the (D/H)_{CH₂DOH} ratios for B1-c and S68N (for B1-bS, both CH₂DOH and CHD₂OH are not detected) are shown in Fig. 4 and agree very well with those derived for other low-mass sources as well as with the high-mass sources.

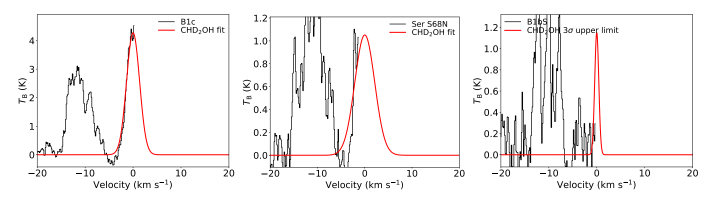


Fig. D.1. Spectral line fits of CHD₂OH $7_{1,2}$ $o_1 - 7_{0,1}$ e_0 ($E_{up} = 68$ K) line for B1-c (left), Serpens S68N (middle), and B1-bS (right). The data corrected for the V_{lsr} are shown in black and the best fit for $T_{ex} = 150$ K is shown in red.

Appendix E: Additional figures

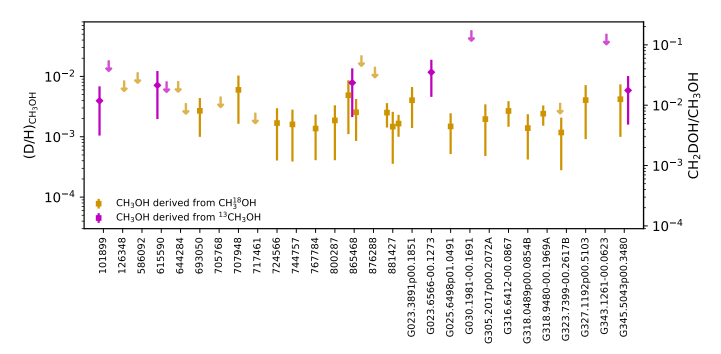


Fig. E.1. (D/H)_{CH₃OH} ratios derived from the $N_{\text{CH}_2\text{DOH}}/N_{\text{CH}_3\text{OH}}$ ratios for the ALMAGAL sources presented in this work, indicating whether $N_{\text{CH}_3\text{OH}}$ was derived from the ^{13}C isotopologue (magenta diamonds) or from the ^{18}O isotopologue (orange squares). Upper limits are presented as arrows

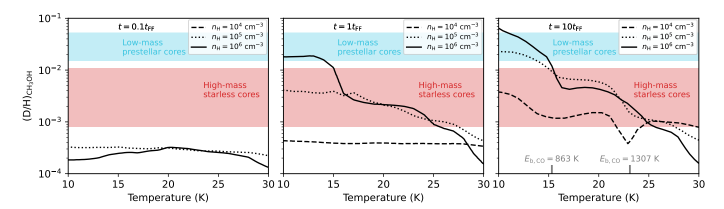


Fig. E.2. Same as Fig. 6 but now showing the average measured $(D/H)_{CH_3OH}$ for low-mass prestellar cores (light blue) and high-mass starless cores (red).