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Protostellar Interferometric Line Survey of the Cygnus X region (PILS-Cygnus)

First results: Observations of CygX-N30[★]

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ABSTRACT

Context. Complex organic molecules (COMs) are commonly detected in and near star-forming regions. However, the dominant process in the release of these COMs from the icy grains – where they predominately form – to the gas phase is still an open question.

Aims. We investigate the origin of COM emission in a high-mass protostellar source, CygX-N30 MM1, through high-angular-resolution interferometric observations over a continuous broad frequency range.

Methods. We used 32 GHz Submillimeter Array observations with continuous frequency coverage from 329 to 361 GHz at an angular resolution of $\sim 1''$ to do a line survey and obtain a chemical inventory of the source. The line emission in the frequency range was used to determine column densities and excitation temperatures for the COMs. We also mapped out the intensity distribution of the different species.

Results. We identified approximately 400 lines that can be attributed to 29 different molecular species and their isotopologues. We find that the molecular peak emission is along a linear gradient, and coincides with the axis of red- and blueshifted H_2CO and CS emission. Chemical differentiation is detected along this gradient, with the O-bearing molecular species peaking towards one component of the system and the N- and S-bearing species peaking towards the other. The chemical gradient is offset from but parallel to the axis through the two continuum sources. The inferred column densities and excitation temperatures are compared to other sources where COMs are abundant. Only one deuterated molecule is detected, HDO, while an upper limit for CH_2DOH is derived, leading to a D/H ratio of $< 0.1\%$.

Conclusions. We conclude that the origin of the observed COM emission is probably a combination of the young stellar sources along with accretion of infalling material onto a disc-like structure surrounding a young protostar and located close to one of the continuum sources. This disc and protostar are associated with the O-bearing molecular species, while the S- and N-bearing species on the other hand are associated with the other continuum core, which is probably a protostar that is slightly more evolved than the other component of the system. The low D/H ratio likely reflects a pre-stellar phase where the COMs formed on the ices at warm temperatures (~ 30 K), where the deuterium fractionation would have been inefficient. The observations and results presented here demonstrate the importance of good frequency coverage and high angular resolution when disentangling the origin of COM emission.

Key words. astrochemistry – stars: protostars – ISM: molecules – ISM: individual objects: W75N(B) – submillimeter: ISM

1. Introduction

Complex organic molecules (COMs; molecules with at least six atoms, of which at least one is carbon; Herbst & van Dishoeck 2009) are found everywhere in and near star-forming regions (e.g. Caselli & Ceccarelli 2012; Ceccarelli et al. 2014; Jørgensen et al. 2020). Our current understanding is that most COMs form

on the surfaces of ice-covered dust grains, either from simple volatile gas species frozen out onto the grains, or from radicals in the ice (Öberg 2016). Near the forming stars, where $T_{\text{dust}} \gtrsim 100$ K, the water-rich ice mantels sublimate off the dust grains, releasing COMs into the gas phase, where they can be directly observed at submillimetre (submm) wavelengths (the so-called hot cores; e.g. Kurtz et al. 2000; Cesaroni et al. 2010; Jørgensen et al. 2016). However, other non-thermal mechanisms can be efficient at desorbing COMs into the gas phase, and have been proposed in the literature. These include sputtering of the

[★] Table 8 is only available at the CDS via anonymous ftp to [cdsarc.u-strasbg.fr](ftp://cdsarc.u-strasbg.fr) (130.79.128.5) or via <http://cdsarc.u-strasbg.fr/viz-bin/cat/J/A+A/655/A86>

ices in shocks, where the shocks are generated by jets and outflows (e.g. Avery & Chiao 1996; Jørgensen et al. 2004; Arce et al. 2008; Sugimura et al. 2011; Lefloch et al. 2017), accretion shocks from the envelope onto the disc (e.g. Podio et al. 2015; Artur de la Villarmois et al. 2018; Csengeri et al. 2018, 2019), explosive events near forming stars (e.g. Orion KL; Zapata et al. 2011; Orozco-Aguilera et al. 2017), and UV irradiation of the outflow cavity walls (e.g. Drozdovskaya et al. 2015). Which of these processes dominate in releasing the molecules from the ice to the gas phase is in many cases unclear, as is the impact on the observed chemistry.

Recently, Belloche et al. (2020) presented observations of a sample of 26 solar-type protostars as part of the CALYPSO survey performed with the Northern Extended Millimeter Array (NOEMA). COM emission from methanol, CH_3OH , the ‘simplest COM’ was detected towards 12 of the sources, with 8 of these sources having detections of at least three COMs. These latter authors found that a canonical hot-corino origin may account for only four of the sources with COM emission, while an accretion-shock origin fits best with two or possibly three sources, and an outflow origin fits three others. It is therefore becoming increasingly clear that the origin of COMs in scenarios other than the canonical hot-corino (and hot cores) are relatively common, and through large surveys like CALYPSO we may get a better understanding of how these COMs form.

The work presented here is part of a large-frequency-range line survey of ten intermediate- to high-mass protostellar sources in the Cygnus X molecular cloud, which forms part of the Great Cygnus Rift in the Galactic plane. Cygnus X is a prolific star-forming region, with many newly formed stars of various masses, and has been the focus of numerous studies (for a full review of the region, see Reipurth & Schneider 2008, and references therein). Given its location in the Perseus Galactic spiral arm, the distance to Cygnus X is uncertain, with values in the literature ranging between 1.3 and 3 kpc (e.g. Campbell et al. 1982; Odenwald & Schwartz 1993; Rygl et al. 2012). We assume the distance of ~ 1.3 kpc obtained by Rygl et al. (2012) from maser parallax measurements. It is home to one of the most massive OB associations known in the Galaxy, Cyg OB2, with hundreds of young O-type stars and perhaps thousands of B-type stars (Knödlseeder 2000). These massive stars provide a huge amount of ionising radiation to the region, making it an ideal target for the study of how the external environment created by these massive stars affects the chemistry of the surrounding star forming regions. To this end, full chemical inventories of a large sample of surrounding protostars are required.

This kind of survey has historically been observationally very expensive, because it requires high angular resolution and many hours of observation time to obtain the wide frequency range needed. However, in recent years it has become much cheaper thanks to the upgraded receivers on the Submillimeter Array (SMA), and its new SWARM¹ correlator. It is now possible to obtain 32 GHz of continuous frequency coverage in a fraction of the time required in the past. Taking advantage of these new capabilities of the SMA, we obtained data from ten newly formed sources surrounding the Cyg OB2 association in order to study how the chemistry of these young stars is affected by the environment in which they form. Together, these observations form the Protostellar Interferometric Line Survey of Cygnus-X (PILS-Cygnus).

The focus of this paper is one of the most studied sources in the Cygnus X cloud, W75N (B) (e.g. Haschick et al. 1981; Persi et al. 2006), or CygX-N30 (N30 hereafter, using the designation by Motte et al. 2007). This source contains three mm continuum cores, MM1, MM2, and MM3, where MM1 is the brightest source and is composed of MM1a and MM1b (e.g. Shepherd 2001; Minh et al. 2010). Three radio continuum sources were also detected towards MM1 (VLA1, VLA2 and VLA3), which have been identified as ultra-compact HII (UC HII) regions or thermal jets (Hunter et al. 1994; Torrelles et al. 1997; Shepherd et al. 2003; Carrasco-González et al. 2010; Rodríguez-Kamenetzky et al. 2020). Shepherd et al. (2004) found that the VLA sources drive high-velocity molecular outflows traced by SiO (2–1 and 1–0) emission. VLA1 and VLA2 were associated with OH, H_2O , and CH_3OH masers (Torrelles et al. 1997; Minier et al. 2001; Hutawarakorn et al. 2002; Fish et al. 2005; Surcis et al. 2009), while VLA3 was found to be associated with one H_2O maser (Shepherd 2001) with a radio jet and SiO (1–0) emission (Carrasco-González et al. 2010). A large-scale CO outflow has been observed and found to originate from MM1 and extending ~ 1 pc in both directions (see Fischer et al. 1985; Hutawarakorn et al. 2002; Shepherd et al. 2003; Gibb et al. 2003; Birks et al. 2006; Surcis et al. 2009, 2011). Through MERLIN² measurements of OH masers, Hutawarakorn et al. (2002) found that N30 hosts a rotating molecular disc with a radius of 3000 AU, a rotation velocity of 6 km s^{-1} , and a mass of $120 M_\odot$. These latter authors found that this disc has a position angle of 155° (measured from north), which is orthogonal to the large-scale molecular outflow seen in CO and H_2 (PA $\sim 65^\circ$). The authors note that the compact cluster of high-velocity OH and H_2O masers coinciding with VLA2 appears to mark the centre of the outflow.

Minh et al. (2010) observed N30 with the SMA in the 215 and 345 GHz spectral windows with 8 GHz frequency coverage, which indicated that a hot core is located at MM1b and is associated with a thermal jet from VLA1. These authors also found a chemical difference between the continuum cores MM1a and MM1b, which they suggest is the result of the evolution of a massive star-forming core. They argue that VLA1 is the heating source of the hot core, which suggests that the region associated with VLA1 is the site of recent star formation. Minh et al. (2010) observed SiO emission at the position of VLA2, which they interpret as a spherical shock driven by the recent star formation at the site of VLA1.

In this paper, we present new SMA observations of N30, providing 32 GHz of continuous frequency coverage in the 345 GHz atmospheric window. This large frequency coverage makes it possible to perform an unbiased survey of the spectroscopic signatures of the different components in the region and establish their molecular inventories. The paper is laid out as follows: a detailed description of the observational setup and data reduction is given in Sect. 2. In Sect. 3, we present the results of our analysis and then discuss them in Sect. 4. We end with a summary and conclusions in Sect. 5.

2. Observations

The SMA observations of N30 form part of the PILS-Cygnus programme (PI: Kristensen, project ID 2017A-S028), which in turn is an extension of the PILS programme (Jørgensen et al. 2016), in which 34 GHz continuous frequency coverage observations of the low-mass protostar, IRAS 16293-2422,

¹ SWARM is an acronym for SMA Wideband Astronomical ROACH2 (second generation Reconfigurable Open Architecture Computing Hardware) Machine.

² Multi-Element Radio Linked Interferometer Network.

obtained from the Atacama Large Millimeter/Submillimeter Array (ALMA) were studied. The ten intermediate- to high-mass protostars of PILS-Cygnus were selected because they are all located in the same molecular cloud structure, and because of their proximity to the Cyg-OB2 association, which provides an opportunity to study the role of the external environment in setting the chemistry of newly formed stars. This paper presents the first results of the PILS-Cygnus programme.

2.1. Calibration

The observations were performed using the SMA in a combination of the compact and extended configurations, that is, with projected baselines ranging from 7 to 211 m. The full survey covered ten sources and every source was observed for approximately an equal amount of time over ten tracks: five tracks in the compact configuration and five tracks in the extended configuration. This was done to ensure approximately equal sensitivity for all observed sources. The compact-configuration tracks were executed between 27 June 2017 and 7 Aug. 2017, while the extended-configuration tracks were observed between 20 Oct. 2017 and 10 Nov. 2017. The number of antennas available in the array was between six and eight.

MWC349A was used as the complex gain calibrator for all observations, while the quasars 3c273, 3c454.3, and 3c84 were used for bandpass calibration depending on the time of the observations. Neptune, Titan, Callisto, and Uranus were used for flux calibration, again depending on the time of the observations. A detailed observing log is presented in Appendix A, which also includes information on the weather at the time of the observations.

The receivers at the SMA were tuned such that they covered the entire frequency range from 329 to 361 GHz continuously. Specifically, the 345 GHz receivers were tuned to cover the spectral range from 329.2 to 337.2 GHz in the lower sideband and the range from 345.2 to 353.2 GHz in the upper sideband. The 400 GHz receivers filled in the 8 GHz gaps, and covered the ranges of 337.2–345.2 GHz and 353.2–361.2 GHz in the lower and upper sidebands, respectively. Each sideband consists of four chunks, and each chunk covers 2 GHz. The SWARM correlator provides a uniform spectral channel size of 140 kHz across the spectrum, corresponding to 0.12 km s^{-1} at 345 GHz. Prior to calibration, all spectral data were rebinned by a factor of four to a spectral resolution of 560 kHz, or 0.48 km s^{-1} at 345 GHz to improve the noise level.

All data were calibrated in CASA 4.7 (Common Astronomy Software Applications, McMullin et al. 2007). Calibration consisted first of flagging the channels at the edges and any anomalously high intensity spikes. The bandpass calibrator was then phase-calibrated with a 30 s solution interval before the complex gains and absolute flux were calibrated.

2.2. Imaging

Self-calibration and imaging were performed using CASA 4.7. Two rounds of phase self-calibration were performed on the continuum, the first with a solution interval of 240 s, followed by a second round with a smaller solution interval of 60 s. This resulted in a reduction in rms level of more than 30%. Only phase self-calibration was performed.

Cleaning and imaging of the data set proceeded along two axes. First, the continuum data were cleaned in order to achieve high angular resolution. This was done by setting the ‘robust’ parameter to -0.5 , where robust = 2 gives natural weighting, and

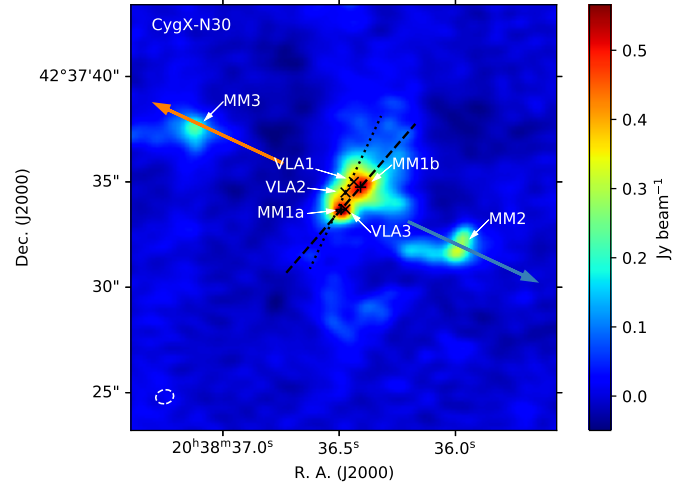


Fig. 1. Observed continuum emission at a frequency of 345 GHz, showing the four continuum sources MM1a, MM1b, MM2, and MM3. The beam is shown in the bottom-left corner, and has dimensions $0''.85 \times 0''.66$, with position angle 12.1° . The VLA radio sources VLA1, VLA2, and VLA3 are labelled and marked with black crosses, with the emission peak positions of MM1a and MM1b shown by plus symbols, and the dashed line representing the axis through these positions. Also shown are the large-scale CO emission (red and blue arrows) and the disc-like structure (dotted line) identified by Hutawarakorn et al. (2002).

robust = -2 gives uniform weighting. This was an affordable solution, because the continuum data had very high signal-to-noise ratio. The resulting beam size of the continuum data is $0''.85 \times 0''.66$ (PA = 12.1°). The continuum rms sensitivity of the final line-free and self-calibrated image (Fig. 1) is $0.008 \text{ Jy beam}^{-1}$.

In a second step, the line data were first continuum-subtracted using line-free channels. This was done by taking a spectrum towards MM1a in the image plane and going through the spectrum by hand to identify all the line-free channels. These frequency ranges were then used throughout the cube, with the continuum subtraction done in the uv -plane. This exercise was then repeated to ensure that all the line-free channels remained line-free. The obtained self-calibration solution was then applied to the line data. To clean the line data, natural weighting was used in order to get the highest possible line sensitivity. For the full data cube, non-interactive cleaning was used with a circular cleaning mask centred at coordinates RA = $20^{\text{h}}38^{\text{m}}36.62^{\text{s}}$ and Dec = $42^\circ37'31''.81$, with a radius of $24''$, and including the MM1, MM2, and MM3 cores. The resulting beam size is $\sim 1''.65 \times 1''.55$ (PA = 11°). The noise level varies across the spectrum from ~ 0.2 – 0.7 Jy beam^{-1} as a function of receiver, as is clearly visible in Fig. 2, particularly around a frequency of 353.2 GHz. The noise level also varies as a function of the Intermediate Frequency (IF) performance of the receiver, as can be seen from the edge effects. A typical value of the rms is $\sim 0.5 \text{ Jy beam}^{-1}$ in 0.48 km s^{-1} channels. The blanked channels in Fig. 2 are high-amplitude edge channels.

3. Results

3.1. Continuum emission

The 345 GHz ($850 \mu\text{m}$) continuum image is shown in Fig. 1. The four previously identified sources are marked as MM1a, MM1b, MM2, and MM3 (e.g. Shepherd 2001; Minh et al. 2010). MM1a

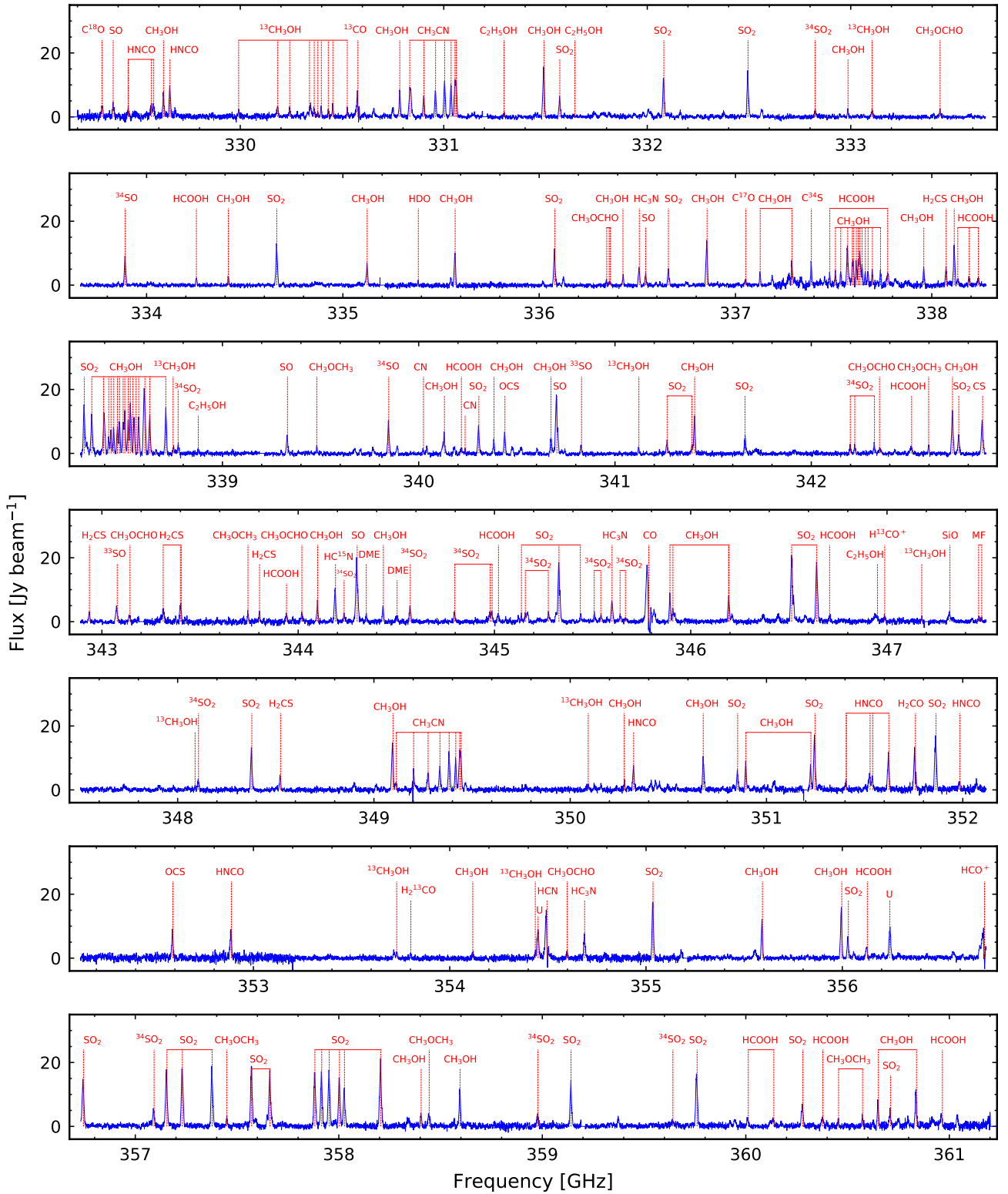


Fig. 2. Spectrum towards N30 at the continuum peak of MM1a. The spectrum shows many molecular lines, with the identified lines marked and labelled. Some abbreviations are used where space is limited; ‘MF’ and ‘DME’ refer to methyl formate (CH_3OCHO) and dimethyl ether (CH_3OCH_3), respectively, while ‘U’ stands for ‘unidentified’.

and MM1b are considerably brighter than MM2 and MM3, peaking at 0.53 and $0.48 \text{ Jy beam}^{-1}$, respectively. All sources show extended emission, with large-scale structure visible to the west of the MM1 region, as well as northwest of MM1b. There is some extended emission to the east of MM2, and some faint

structure just visible to the south of MM1. As with previous studies (e.g. Minh et al. 2010), we detect no COM emission from the continuum cores MM2 and MM3, and this paper therefore focuses on emission towards MM1a and MM1b. The positions of the peak continuum emission for MM1a and MM1b were

Table 1. Properties of the N30-MM1 continuum cores.

Name	Right ascension	Declination	$S_{850\ \mu\text{m}}$ (Jy beam ⁻¹)
MM1a	20 ^h 38 ^m 36 ^s .51	42° 37′ 33″.48	0.53
MM1b	20 ^h 38 ^m 36 ^s .42	42° 37′ 34″.58	0.48

derived using 2D Gaussian fits, and are shown in Table 1 and marked with plus symbols in Fig. 1, with the dashed line representing the axis going through these two positions. MM1a and MM1b are separated by $\sim 1''$, or ~ 1300 AU, at a distance of 1.3 kpc.

In addition to the 345 GHz continuum cores, Fig. 1 also shows the positions of the three radio continuum sources VLA1, VLA2, and VLA3, which were identified previously (Hunter et al. 1994; Torrelles et al. 1997; Shepherd et al. 2003) using mm and centimetre (cm) observations from the Very Large Array (VLA) telescope. VLA1 is located to the west and slightly to the north of the MM1b submm peak, while VLA2 is to the north of the MM1a peak (southwest of MM1b). VLA3 is located very close to but slightly to the east of MM1a (see also Rodríguez-Kamenetzky et al. 2020, for recent high-resolution VLA observations of the region). The red and blue arrows represent the large-scale CO emission (PA $\sim 65^\circ$ from north) centred at VLA2, with the dotted line representing the disc-like structure (PA $\sim 155^\circ$) traced by OH masers, as identified by Hutawarakorn et al. (2002).

To check whether the dust is optically thick or not, the first step is to calculate the dust mass. Here, this is only done towards the MM1a source, which is the brightest continuum source in the field. The total mass (gas + dust) can be calculated using the following relation (e.g. Artur de la Villarmois et al. 2018):

$$M = \frac{S_\nu d^2}{\kappa_\nu B_\nu(T)}, \quad (1)$$

where $S_\nu = 0.53$ Jy beam⁻¹ is the peak intensity of MM1a, $d = 1.3$ kpc, and $B_\nu(T)$ is the Planck function at the specific frequency and temperature. κ_ν is the dust opacity (at $\nu = 345$ GHz, $\kappa_\nu = 0.0175$ cm² per gram of gas for a gas-to-dust ratio of 100; Ossenkopf & Henning 1994). For a beam size of $0''.85 \times 0''.66$, and taking $T = 30$ K, we get $M = 3.0 M_\odot$ of gas and dust. If this mass is spread evenly over the beam, this corresponds to an H₂ column density of 5.3×10^{24} cm⁻², where a mean molecular weight of $\mu = 2.8$ was used to account for Helium (Kauffmann et al. 2008). This may be converted to optical depth using the following expression (e.g. Schöier et al. 2002):

$$\tau_\nu = \kappa_\nu \mu m_H N_{H_2}. \quad (2)$$

We note that Schöier et al. (2002) includes a dust-to-gas ratio, $\delta = 0.01$, and κ is their dust opacity (dust only). As we have folded the dust-to-gas ratio into κ_ν above (gas + dust), this parameter is not needed in the expression for the dust optical depth. Furthermore, we correct a typo in Schöier et al. (2002) and use μm_H instead of m_{H_2} , again with $\mu = 2.8$. This yields a dust opacity of 0.43. We conclude that the dust emission is at least marginally optically thin for a gas-to-dust ratio of 100.

3.2. Molecular line emission

Figure 2 shows a spectrum from the data taken at the position of peak continuum emission of MM1a. The 32 GHz continuous

frequency coverage of the data (329–361 GHz) included approximately 400 molecular lines detected above 3σ towards the MM1 source, or a line temperature sensitivity of ~ 10 K. Line identification was done by covering the points as described in Snyder et al. (2005) and using the software package CASSIS³ (Centre d’Analyse Scientifique de Spectres Instrumentaux et Synthétiques; Vastel et al. 2015), in which the CDMS⁴ (Müller et al. 2001, 2005; Endres et al. 2016) and JPL⁵ (Pickett et al. 1998) molecular spectroscopy databases were used to construct a synthetic spectrum. Previously identified molecules were added to the synthetic spectrum first, which was then compared to the observed spectrum in order to find and identify the remaining lines above 3σ . From the observed lines, we identified 29 different molecules and their isotopologues, including five COMs, (CH₃OH, C₂H₅OH, CH₃OCH₃, CH₃OCHO, and CH₃CN).

The molecular transitions detected at $>3\sigma$ are listed below, with the oxygen-bearing species in Table 2, sulphur-bearing species in Table 3, and nitrogen-bearing species shown in Table 4, together with SiO, HDO, CO, and HCO⁺. There are approximately 40 lines that remain unidentified, including two bright lines (at frequencies not corrected for the systemic velocity of 354.44896 GHz and 356.24131 GHz, and labelled ‘U’ in Fig. 2). A full list of detected molecules above 3σ are listed in Table 8.

3.3. Integrated molecular intensity maps

The molecular peak positions were found by making integrated intensity emission maps of each molecule. Where a molecule had more than one line, the lines were stacked using a $1/\sigma^2$ weighting scheme – where σ is the RMS measured on an emission-free region for each channel map – in order to increase the intensity and determine an average peak position of the molecule. Only unblended lines were used.

In the cases where red- and blueshifted components are present, the components were separated with integrated maps made for each component. A map of blue- and redshifted H₂CO emission is shown in Fig. 3 as an example (frequency = 351.77 GHz, $J = 5_{1,5}-4_{1,4}$, and $E_{\text{up}} = 62.45$ K). In this figure, the continuum emission is depicted in colour, with the red- and blueshifted integrated intensity maps shown in red and blue contours, respectively.

The maximum intensity for the components were found at positions offset from the MM1a continuum peak: $(-0''.83, 1''.72)$ for the blueshifted components and $(-0''.03, 0''.52)$ for the redshifted ones. The line plots for the blue- and redshifted components are shown in Figs. 4 and 5, respectively.

As can be seen from Fig. 3, the blue- and redshifted components of H₂CO emission falls on an axis – marked with a dotted line – that runs parallel to the axis drawn between the positions of the MM1a and MM1b peak continuum emission marked with plus signs and represented with a dashed line. The blue contours are more collimated along the axis, but extend further out than the red contours, while the red contours have some extended emission to the west of MM1b, and north of the MM2 core. Similarly, the red- and blueshifted 342.88 GHz CS $J = 7-6$ transition is shown in contours in Fig. 6, with the line plots for the two components shown in Figs. 7 and 8. Other molecules that exhibit red- and blueshifted components are CO, ¹³CO, HCN, and HCO⁺.

³ <http://cassis.irap.omp.eu>

⁴ <https://cdms.astro.uni-koeln.de/>

⁵ <http://spec.jpl.nasa.gov/>

Table 2. Gaussian fits for molecular transitions of COMs, including t-HCOOH and H₂CO.

Molecule	Frequency (GHz)	Transition	Position (",")	$I_{\text{peak}}^{(a)}$ (Jy beam ⁻¹)	$I_{\text{int}}^{(a)}$ (Jy beam ⁻¹ km s ⁻¹)	$FWHM^{(a)}$ (km s ⁻¹)	$v_{\text{source}}^{(a)}$ (km s ⁻¹)
A-CH ₃ OH	331.5023	11 _{1,-0} –11 _{0,+0}	(−0.31, 0.70)	16.0	123.0	8.5	9.9
	336.8651	12 _{1,-0} –12 _{0,+0}	(−0.33, 0.82)	14.9	112.4	8.2	10.2
	338.4087	7 _{0,+0} –6 _{0,+0}	(−0.48, 0.85)	14.7	124.6	10.4	9.9
	341.4156	7 _{1,-0} –6 _{1,-0}	(−0.29, 0.86)	12.6	87.7	8.0	8.7
	342.7298	13 _{1,-0} –13 _{0,+0}	(−0.27, 0.75)	14.9	109.5	8.3	9.1
	349.1070	14 _{1,-0} –11 _{0,+0}	(−0.40, 0.76)	15.7	120.4	8.3	11.0
	355.6029	13 _{0,+0} –11 _{1,+0}	(−0.41, 1.03)	13.9	99.5	7.5	10.4
	356.0072	15 _{1,-0} –15 _{0,+0}	(−0.37, 0.82)	17.1	134.4	8.5	10.8
	360.6616	3 _{1,+1} –4 _{2,+1}	(−0.45, 0.98)	10.6	76.1	7.4	9.5
E-CH ₃ OH	338.1244	7 _{0,0} –6 _{0,0}	(−0.45, 0.87)	14.8	102.8	7.4	9.8
	338.3446	7 _{−1,0} –6 _{−1,0}	(−0.50, 0.90)	14.5	104.9	8.2	9.8
	360.8489	11 _{0,0} –10 _{1,0}	(−0.41, 0.93)	13.6	100.9	7.8	10.0
¹³ CH ₃ OH	330.0018	7 _{0,7,0} –6 _{0,6,0}	(−0.59, 0.99)	3.5	18.8	5.8	7.9
	330.2528	7 _{0,7,+0} –6 _{0,6,+0}	(−0.55, 1.07)	3.8	24.6	8.0	8.3
	330.4424	7 _{1,6,0} –6 _{1,5,0}	(−0.51, 0.93)	3.5	16.7	5.5	8.0
	330.4639	11 _{1,10,-0} –11 _{0,11,+0}	(−0.52, 0.94)	6.1	32.1	6.1	7.7
	330.5352	7 _{2,5,0} –6 _{2,4,0}	(−0.53, 1.02)	4.9	26.6	6.1	7.6
	333.1148	7 _{1,6,-0} –6 _{1,5,-0}	(−0.61, 0.98)	3.5	22.6	6.6	8.8
	335.5602	12 _{1,11,-0} –12 _{0,12,+0}	(−0.57, 1.05)	4.0	22.8	5.8	9.2
C ₂ H ₅ OH	338.8862	15 _{7,8,2} –15 _{6,9,2}	(−0.94, 1.45)	1.9	11.1	8.4	7.6
	339.3126	12 _{7,5,2} –12 _{6,6,2}	(−1.09, 1.67)	1.1	7.8	7.6	9.1
	339.3984	11 _{7,4,2} –11 _{6,5,2}	(−1.12, 1.63)	1.0	6.4	7.8	8.4
CH ₃ OCH ₃	339.4916	19 _{1,18,5} –18 _{2,17,5}	(−0.54, 1.04)	3.5	18.6	5.2	9.4
	340.6126	10 _{3,7,1} –9 _{2,8,1}	(−0.89, 1.21)	2.9	16.2	6.6	7.7
	342.6080	19 _{0,19,0} –18 _{1,18,0}	(−0.57, 1.22)	4.7	27.7	5.7	9.4
	344.3578	19 _{1,19,3} –18 _{0,18,3}	(−0.64, 1.15)	4.4	27.3	6.6	8.8
	357.4602	18 _{2,17,1} –17 _{1,16,1}	(−0.69, 1.08)	4.8	28.3	7.0	8.0
	358.4541	5 _{5,1,1} –4 _{4,1,1}	(−0.76, 1.17)	7.3	63.0	10.6	10.2
	360.4660	20 _{1,19,0} –19 _{2,18,0}	(−0.79, 1.22)	4.0	25.9	6.1	10.9
	360.5847	20 _{0,20,0} –19 _{1,19,0}	(−0.80, 1.23)	5.3	35.8	6.9	10.1
CH ₃ OCHO	333.4494	31 _{1,31,0} –30 _{0,30,0}	(−0.69, 1.14)	4.0	23.5	6.0	8.9
	336.3514	27 _{6,22,1} –26 _{6,21,1}	(−0.82, 1.24)	2.2	15.6	7.0	10.3
	344.0296	19 _{1,18,5} –18 _{2,17,5}	(−0.52, 1.08)	4.4	29.4	7.1	10.1
	354.6081	19 _{1,18,5} –18 _{2,17,5}	(−0.65, 1.23)	4.0	24.4	6.7	9.5
t-HCOOH	334.2658	15 _{2,14} –14 _{2,13}	(−0.03, 0.46)	2.5	15.8	6.9	10.2
	343.9523	15 _{1,14} –14 _{1,13}	(0.03, 0.37)	2.4	17.1	7.8	11.2
H ₂ CO ^(b)	351.7686	5 _{1,5} –4 _{1,4}	(−0.03, 0.52)	13.1	88.0	6.3	13.9
		5 _{1,5} –4 _{1,4}	(−0.83, 1.72)	11.0	47.1	3.7	6.5

Notes. The transitions listed here, as well as the transitions listed in Tables 3 and 4, were relatively isolated and unblended, and were used to make the emission maps shown in Fig. 11. ^(a)Typical uncertainties on the Gaussian centroid fits are (0'2, 0'2) for the position, 0.3 Jy beam⁻¹ for I_{peak} , Jy beam⁻¹ km s⁻¹ for I_{int} , and 0.4 km s⁻¹ for the FWHM and v_{source} . ^(b)The two entries for H₂CO are red- and blueshifted lines, respectively. See also entries for CS.

In order to investigate this large-scale velocity gradient, Figs. 9 and 10 show position–velocity maps of the H₂CO and CS emission, respectively. The high-velocity emission of H₂CO and CS is close to the centre (at 0'' offset, between the red- and blueshifted emission peaks), whereas the emission at a larger offset from the centre shows a velocity closer to the systemic velocity of the protostar. This could trace an infall motion coupled to a rotation pattern, which might be due to a disc-like structure (e.g. Zhu et al. 2011). A comprehensive study of the kinematics of the system is beyond the scope of this paper, but

position–velocity maps of a selection of other transitions are shown in Appendix C.

The panels in Fig. 11 show the molecular peaks of each molecule; these peaks are concentrated along the axis of the red- and blueshifted components of H₂CO and CS emission. Each panel shows a contour map of the integrated molecular line emission of a different molecule. In the case of molecules with more than one unblended line, such as for example CH₃OH, the images show stacked emission for better S/N. The peak for each molecule was found using 2D Gaussian fits to the integrated

Table 3. Gaussian fits for molecular transitions of S-bearing species.

Molecule	Frequency (GHz)	Transition	Position ($''$, $''$)	I_{peak} (Jy beam $^{-1}$)	I_{int} (Jy beam $^{-1}$ km s $^{-1}$)	$FWHM$ (km s $^{-1}$)	v_{source} (km s $^{-1}$)
CS	342.8828	7 ₀ –6 ₀	(0.17, 0.72)	11.1	69.7	8.5	9.7
	342.8828	7 ₀ –6 ₀	(−1.03, 1.92)	9.2	47.3	4.2	7.9
C ³⁴ S	337.3965	7 ₀ –6 ₀	(−0.54, 1.08)	8.3	49.1	6.0	9.0
SO	340.7141	8 ₇ –6 ₆	(0.13, 0.24)	16.3	84.9	9.5	5.4
	334.4311	8 ₈ –7 ₇	(0.14, 0.25)	17.7	98.5	10.6	5.2
	346.5281	8 ₉ –7 ₈	(0.06, 0.25)	19.4	140.0	11.1	6.7
³⁴ SO	333.9010	8 ₇ –7 ₆	(−0.07, 0.22)	8.1	57.4	7.5	9.9
	337.5801	8 ₈ –7 ₇	(−0.22, 0.29)	10.7	80.9	9.0	8.7
	339.8573	8 ₉ –7 ₈	(−0.16, 0.18)	9.0	64.1	7.5	10.5
SO ₂	332.0914	21 _{2,20} –21 _{1,21}	(−0.14, 0.17)	11.4	86.6	8.0	10.8
	332.5052	4 _{3,1} –3 _{2,2}	(−0.06, 0.30)	12.9	95.7	8.3	9.9
	334.6733	8 _{2,6} –7 _{1,7}	(−0.09, 0.23)	12.0	85.6	7.8	9.5
	336.0892	23 _{3,21} –23 _{2,22}	(−0.15, 0.29)	10.1	76.5	8.2	10.9
	336.6695	16 _{7,9} –17 _{6,12}	(0.03, 0.14)	4.8	35.2	7.6	10.8
	338.3060	18 _{4,14} –18 _{3,15}	(−0.06, 0.23)	14.3	107.0	8.3	10.0
	340.3164	28 _{2,26} –28 _{1,27}	(−0.05, 0.27)	8.0	58.5	7.9	10.6
	345.3385	13 _{2,12} –12 _{1,11}	(−0.15, 0.33)	16.7	137.2	10.6	10.4
	346.6522	19 _{1,19} –18 _{0,18}	(−0.09, 0.27)	17.4	133.7	9.1	9.4
	350.8628	10 _{6,4} –11 _{5,7}	(0.01, 0.07)	5.6	41.7	8.1	9.8
	351.8739	14 _{4,10} –14 _{3,11}	(−0.11, 0.19)	15.2	116.2	8.7	9.9
	355.0455	12 _{4,8} –12 _{3,9}	(−0.09, 0.27)	16.2	122.4	8.5	9.7
	357.1654	13 _{4,10} –13 _{3,11}	(−0.21, 0.37)	16.1	124.4	9.2	11.6
	357.2412	15 _{4,12} –15 _{3,13}	(−0.19, 0.29)	17.0	129.7	8.7	10.3
	357.3876	11 _{4,8} –11 _{3,9}	(−0.15, 0.27)	16.4	129.2	9.1	10.2
	357.5814	8 _{4,4} –8 _{3,5}	(−0.14, 0.22)	17.0	131.8	9.0	10.2
	357.8924	7 _{4,4} –7 _{3,5}	(−0.11, 0.27)	15.7	123.6	9.2	10.2
	357.9258	6 _{4,2} –6 _{3,3}	(−0.15, 0.29)	15.6	121.3	9.1	10.5
	357.9629	17 _{4,14} –17 _{3,15}	(−0.09, 0.26)	15.7	126.6	9.4	10.0
	358.0131	5 _{4,2} –5 _{3,3}	(−0.27, 0.37)	13.6	105.0	8.9	10.1
	358.2156	20 _{0,20} –19 _{1,19}	(−0.17, 0.25)	19.0	150.8	9.5	10.4
	359.7707	19 _{4,16} –19 _{3,17}	(−0.16, 0.29)	15.5	113.7	8.9	12.0
³⁴ SO ₂	342.2089	5 _{3,3} –4 _{2,2}	(−0.03, 0.37)	2.4	15.4	7.2	9.7
	342.3320	12 _{4,8} –12 _{3,9}	(0.08, 0.29)	2.8	19.7	7.2	10.5
	344.2453	10 _{4,6} –10 _{3,7}	(−0.02, 0.25)	2.3	17.7	9.1	10.5
	344.5810	19 _{1,19} –18 _{0,18}	(−0.06, 0.20)	4.5	35.3	8.3	11.0
	344.8079	13 _{4,10} –13 _{3,11}	(−0.05, 0.21)	2.7	20.0	8.3	11.0
OCS	340.4493	28–27	(−0.45, 0.79)	8.2	55.4	6.8	9.8
	352.5996	29–28	(−0.48, 0.70)	9.4	70.8	7.7	10.2
OC ³⁴ S	332.1297	28–27	(−1.04, 1.44)	1.1	6.0	7.8	10.9
	343.9833	29–28	(−1.04, 1.49)	2.0	11.7	5.4	9.6
H ₂ CS	338.0832	10 _{1,10} –9 _{1,9}	(−0.67, 1.16)	7.7	47.4	6.8	8.4
	342.9464	10 _{0,10} –9 _{0,9}	(−0.72, 1.22)	4.4	26.6	6.6	8.3
	343.4141	10 _{3,7} –9 _{3,6}	(−0.64, 1.00)	4.3	23.0	5.1	9.0
	343.8132	10 _{2,8} –9 _{2,7}	(−0.73, 1.23)	7.1	46.1	6.8	9.7

Notes. Uncertainties are as in Table 2.

(and again stacked where more than one unblended line was detected) molecular line emission, and is indicated with a plus symbol of the same colour as the contour for that molecule. To verify these fit results for the peak positions, an independent fit of the strongest unblended CH₃OH and SO₂ lines was performed in uv -space prior to cleaning the data. These fits confirm the results from the image plane, and typical differences are less than 0''.2, the adopted pixel size, and one-fifth of the beam. We

therefore adopt a typical uncertainty on the peak locations as 0''.2 throughout. However, we note that the formal uncertainties on the peak locations from the fits in the uv -plane and image plane, respectively, yield uncertainties of typically 0''.01–0''.02.

The full width at half maximum (FWHM) and source velocities (v_{source}) of the lines were determined by making 1D Gaussian fits to the respective lines. The detailed values obtained for each line are shown in Table 2 for the oxygen-, Table 3 for the

Table 4. Gaussian fits for molecular transitions of N-bearing species, CO, HCO⁺, SiO and HDO.

Molecule	Frequency (GHz)	Transition	Position (",")	I_{peak} (Jy beam ⁻¹)	I_{int} (Jy beam ⁻¹ km s ⁻¹)	$FWHM$ (km s ⁻¹)	v_{source} (km s ⁻¹)
N-bearing species							
CH ₃ CN	330.7603	18 ₇ –17 ₇	(−0.22, 0.50)	6.1	46.2	7.4	10.0
	330.9126	18 ₅ –17 ₅	(−0.18, 0.47)	7.8	56.2	7.1	9.5
	330.9698	18 ₄ –17 ₄	(−0.22, 0.51)	11.4	82.3	7.5	9.2
	331.0143	18 ₃ –17 ₃	(−0.24, 0.55)	10.0	72.6	7.2	9.2
	331.0461	18 ₂ –17 ₂	(−0.35, 0.57)	7.5	55.3	7.4	9.5
	349.3933	19 ₃ –18 ₃	(−0.20, 0.56)	11.7	88.8	8.1	9.7
	349.4268	19 ₂ –18 ₂	(−0.20, 0.60)	9.8	73.0	7.7	10.1
HNCO	329.6644	15 _{0,15} –14 _{0,14}	(−0.14, 0.42)	8.8	66.9	8.3	9.2
	350.3330	16 _{1,16} –15 _{1,15}	(−0.05, 0.35)	7.6	61.4	8.4	10.2
HCN ^(a)	354.5055	4–3	(0.17, 0.52)	17.8	127.2	6.3	14.2
HC ¹⁵ N	344.2001	4–3	(−0.19, 0.59)	10.2	78.7	8.6	9.9
HC ₃ N	336.5201	37–36	(−0.14, 0.53)	5.6	44.2	7.9	10.7
	345.6090	38–37	(−0.15, 0.47)	6.5	51.8	8.6	10.1
	354.6975	39–38	(−0.18, 0.39)	6.7	53.8	8.7	10.6
CN	340.2478	3 _{0,3.5} –2 _{0,2.5}	(−0.49, 1.41)	1.4	11.3	9.8	9.4
SiO and HDO							
SiO	347.3306	8 ₀ –7 ₀	(−0.07, 0.47)	2.8	33.6	11.7	11.2
HDO ^(b)	335.39550	3 _{3,1} –4 _{2,2}	(−0.43, 0.92)	1.6	11.5	2.1	10.8
CO and HCO ⁺							
CO ^(a)	345.7960	3–2	(0.35, 0.68)	18.2	156.0	10.1	11.6
¹³ CO	330.5880	3–2	(0.57, −0.12)	10.7	51.9	4.5	10.6
C ¹⁸ O	329.3305	3–2	(−0.64, 0.37)	3.5	25.2	8.2	9.1
H ¹³ CO ⁺	346.9983	4–3	(−1.45, 0.50)	3.74	–	4.31	9.20

Notes. Uncertainties are as in Table 2. ^(a)For the HCN and CO lines, only the brightest, redshifted components are listed. ^(b)HDO emission is extended, so the peak position is of the maximum flux position, not derived from a 2D Gaussian fit.

sulphur-, and Table 4 for the nitrogen-bearing species, with this table also showing SiO, HDO, CO, and HCO⁺.

3.4. Molecular gradient

We find that all molecules that do not show significant extended emission peak along a linear gradient coinciding with the axis marked by the red- and blueshifted components of the H₂CO and CS emission. This axis runs parallel to but is offset from the axis drawn between the continuum peaks of MM1a and MM1b, as shown in Fig. 12. To investigate whether or not this observed molecular gradient is the result of a difference in upper-level energy of the molecular transitions, that is, a temperature gradient, the peak positions of different transitions of CH₃OH and SO₂ are represented in Figs. 13 and 14, where the upper-level energies are represented in different colours. As can be seen from the plot, there does not seem to be any pattern, and the positions of the respective lines seem to be distributed randomly. More specifically, the peaks are distributed within less than one-third of the beam, that is, the spread is over ~0'4, whereas the beam size is ~1'5. A similar exercise was performed for the critical densities (n_{crit}), where $n_{\text{crit}} = A_{ij}/K_{ij}$, with A_{ij} the Einstein A coefficient for spontaneous emission, and K_{ij} the rate coefficient

of the transition from the upper level i to the lower level j , for temperatures of 100 K (rate coefficients were obtained from the Leiden Atomic and Molecular Database; Schöier et al. 2005)⁶. Figures 15 and 16 again show the peak positions of the different transitions of CH₃OH and SO₂, with the critical densities of different transitions represented in colours. As with the upper-level energies, there does not seem to be any density gradient, and the peaks are localised within a fraction of a beam (0'4–0'6 vs. the 1'5 beam), which leads us to conclude that the observed gradient is not an excitation gradient.

Figure 12 illustrates the observed molecular gradient, where the molecular peak emission is represented in three groups. The O-bearing COMs peak closer to MM1b, between VLA1 and VLA2 (positions 1 and 2), with CH₃OH peaking close to VLA2, and the more complex species (C₂H₅OH, CH₃OCHO, and CH₃OCH₃) peaking closer to VLA1. The exceptions here are the S-bearing species, CS, OCS, and H₂CS, which peak between VLA1 and VLA2, while the other S-bearing species peak between VLA2 and VLA3 (position 3). This is also where the N-bearing species peak. Moreover, HCOOH peaks close to the VLA3/MM1a positions, but shows more extended emission

⁶ <https://home.strw.leidenuniv.nl/~moldata/>

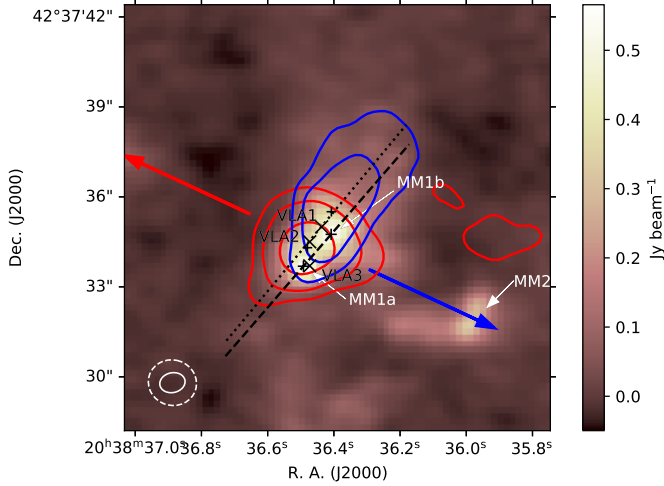


Fig. 3. Continuum image of N30, with the red and blueshifted 351.77 GHz H_2CO , $J = 5_{1,5}-4_{1,4}$, transition shown in contours. The blueshifted component shows more concentrated emission along the outflow axis (represented with a dotted line, running nearly parallel to the axis through the continuum peaks, represented with a dashed line), while the redshifted emission is slightly more extended perpendicular to the outflow axis, with some emission to the north of the MM2 continuum core. The velocity ranges extend to $\pm 14.25 \text{ km s}^{-1}$ from the systemic velocity of 9.5 km s^{-1} . Beam sizes are shown in the bottom left corner, and are $0''.85 \times 0''.66$, with $\text{PA} = 12.1^\circ$, for the continuum, and $1''.61 \times 1''.51$, $\text{PA} = 162.24^\circ$, for the molecular emission. Contour levels are at 3σ , 6σ , and 12σ . The blue and red arrows represent the large-scale CO emission.

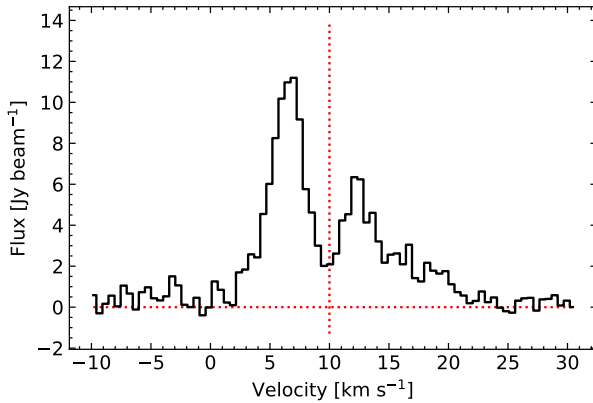


Fig. 4. Blueshifted peak of the 351.77 GHz H_2CO , $J = 5_{1,5}-4_{1,4}$ line, which peaks at an offset of $(-0.075, 1.72)$ arcseconds from the MM1a continuum peak (see Table 2).

in the northwest direction, along the blueshifted outflow axis (see Fig. 11).

3.5. Other molecules

The 347.33 GHz SiO , $J = 8-7$, transition has a weak blueshifted component at -6 km s^{-1} (or perhaps a neighbouring unidentified line, see Fig. 17), which was included in the integrated line map, with the velocity range of the integrated map from -10 km s^{-1} to 20 km s^{-1} . The source velocity and FWHM listed in Table 4 are for the component centred around 10 km s^{-1} . The molecular peak for SiO is at position 3 in Fig. 12, together with the S- and N-bearing species.

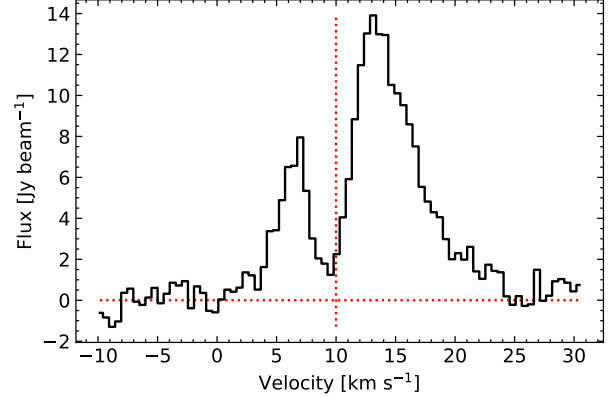


Fig. 5. Redshifted peak of the 351.77 GHz H_2CO , $J = 5_{1,5}-4_{1,4}$ line, which peaks at an offset of $(-0.003, 0.52)$ arcseconds from the MM1a continuum peak (see Table 2).

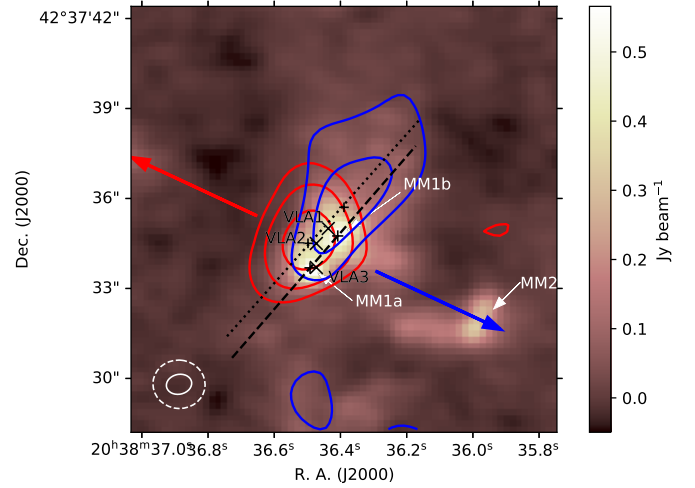


Fig. 6. Continuum image of N30, with the red- and blueshifted 342.88 GHz CS line transition $J = 7-6$ shown in contours. As with the H_2CO line, CS shows red- and blueshifted emission along the same outflow axis. The blueshifted emission shows some extended emission further out to the northeast, while the redshifted emission is slightly more concentrated than for H_2CO . Here, the velocity ranges extend to $\pm 15.6 \text{ km s}^{-1}$ from the systemic velocity of 8.75 km s^{-1} . Beam sizes and contour levels are as in Fig. 3.

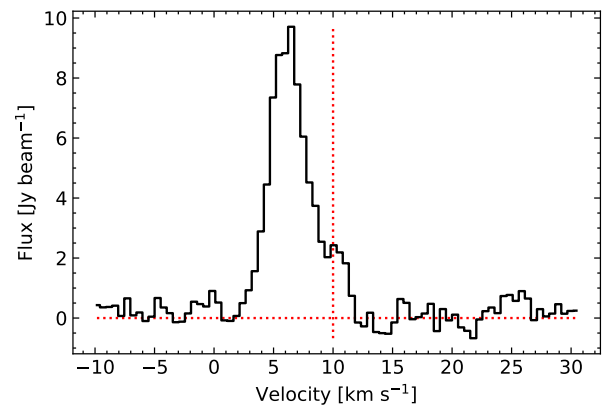


Fig. 7. Blueshifted peak of the 342.88 GHz CS $J = 7-6$ line, which peaks at an offset of $(-0.094, 1.92)$ arcseconds from the MM1a continuum peak (see Table 3).

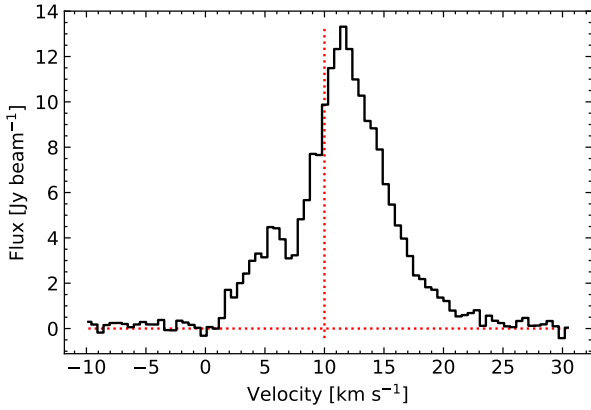


Fig. 8. Redshifted peak of the 342.88 GHz CS $J = 7-6$ line, which peaks at an offset of (0.015, 0.72) arcseconds from the MM1a continuum peak (see Table 3).

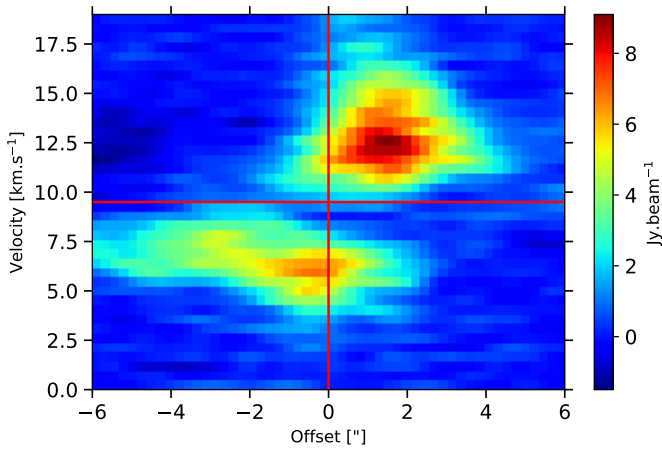


Fig. 9. Position–velocity map of the 351.77 GHz H₂CO, $J = 5_{1,5}-4_{1,4}$ line. The high-velocity emission is located close to the position of zero offset (located between the red- and blueshifted emission peaks), whereas the emission at larger offsets from the centre has velocity closer to the systemic velocity.

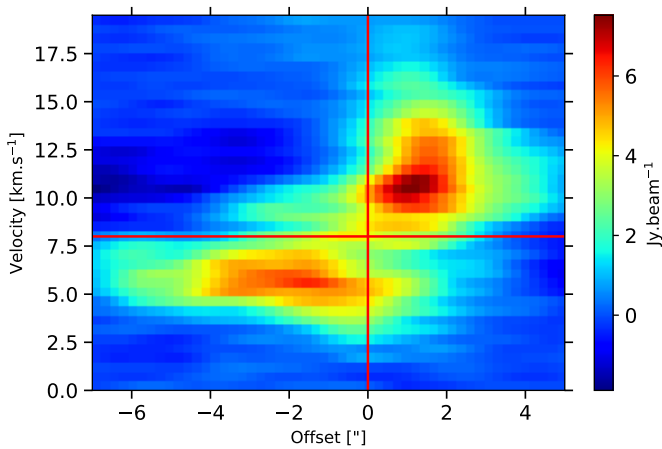


Fig. 10. Position–velocity map of the 342.88 GHz CS $J = 7-6$ line. As with the H₂CO map, the emission at larger velocity is concentrated close to the centre of the emission (between the red- and blueshifted peaks), whereas the emission close to the systemic velocity extends out to larger spacial scales.

Only one deuterated molecule is detected, the 335.395 GHz HDO, $J = 3_{3,1}-4_{2,2}$ transition. This is shown in contours over the continuum in Fig. 18. HDO shows extended emission, also following the red- and blueshifted H₂CO and CS axes, but with some elongated structure to the north of VLA1 and MM1b. The line peaks at 2.39 Jy beam⁻¹, or about 6σ , and with the peak position located between the VLA1 and VLA2 peaks.

3.6. Column densities

As we observe such a strong chemical gradient in the positions of the molecular peak emission, we derived column densities for the respective molecules at the three positions shown in Fig. 12. A synthetic spectrum was constructed with CASSIS using only line emission, assuming local thermodynamic equilibrium (LTE). The inbuilt regular grid function was used to cover a large parameter space and compute the reduced χ^2 minimum in order to determine the best-fit spectral model. Four parameters were used as variables for fitting of the observed spectra. These were the source velocity (v_{source}), FWHM, the column density (N), and excitation temperature (T_{ex}). Estimates for the FWHM and v_{source} were first obtained by looking at a few individual lines and fitting Gaussian profiles to these lines. These estimates were then used with a parameter space of 4 km s⁻¹ around the lines, in steps of 0.5 km s⁻¹, which is equivalent to the channel width. The initial parameter space for N was $N_{\text{min}} = 10^{14}$ and $N_{\text{max}} = 10^{19}$, with ten steps in the range, and for T_{ex} the range was 90 to 300 K, also with ten steps in this range. A few iterations were then run to decrease the ranges, but keeping the step size larger than the uncertainties. The source size was assumed to be 1.5'', or slightly larger than the beam size. Only line data were used in the modelling process. The values obtained for each molecule are shown in Tables 5, 6, and 7. Table 5 shows the values obtained from fits to the spectrum taken at position 1 (see Fig. 12), where the O-bearing species peak around VLA1. Table 6 in turn shows values obtained at peak position 2, between VLA1 and VLA2. Finally, Table 7 shows the values obtained at position 3, where the N- and S-bearing species peak (between VLA2 and VLA3). The uncertainties for N and T_{ex} are mainly a result of how well the spectral model fits the data. This in turn depends on our assumptions of LTE (if the level populations are not in LTE, this will introduce a systematic error), optically thin emission (if emission is optically thick, column densities will be underestimated), and the line shape (we assume Gaussian line profiles). For species with many optically thick lines, this becomes especially problematic because fewer lines can be used in the fit. The optical depth of each line was checked, and only lines with $\tau \lesssim 0.8$ were used in the modelling. The uncertainties on N and T_{ex} were obtained by fixing N (or T_{ex}), and then changing T_{ex} (or N) until we were able to observe the impact of the change on the resulting fit, following the method outlined in Calcutt et al. (2018). Figures of each of the fitted lines for the molecules at position 2 are shown in Appendix D, with the synthetic spectra shown in red and over-plotted on the observed spectra in black.

In the case of molecules with many optically thick lines, such as CH₃OH and SO₂, the column densities were obtained from optically thin isotopologue emission. The isotopologue column density was then multiplied by the ISM isotope ratio. As an example, the modelled column density of ¹³CH₃OH was multiplied by the ¹²C/¹³C ratio, which we took to be ~ 77 (Wilson & Rood 1994). Similarly, column densities for optically thick sulphur-bearing molecular lines were obtained using the ISM isotopologue ratio ³²S/³⁴S ~ 22 (Wilson 1999). In the case of

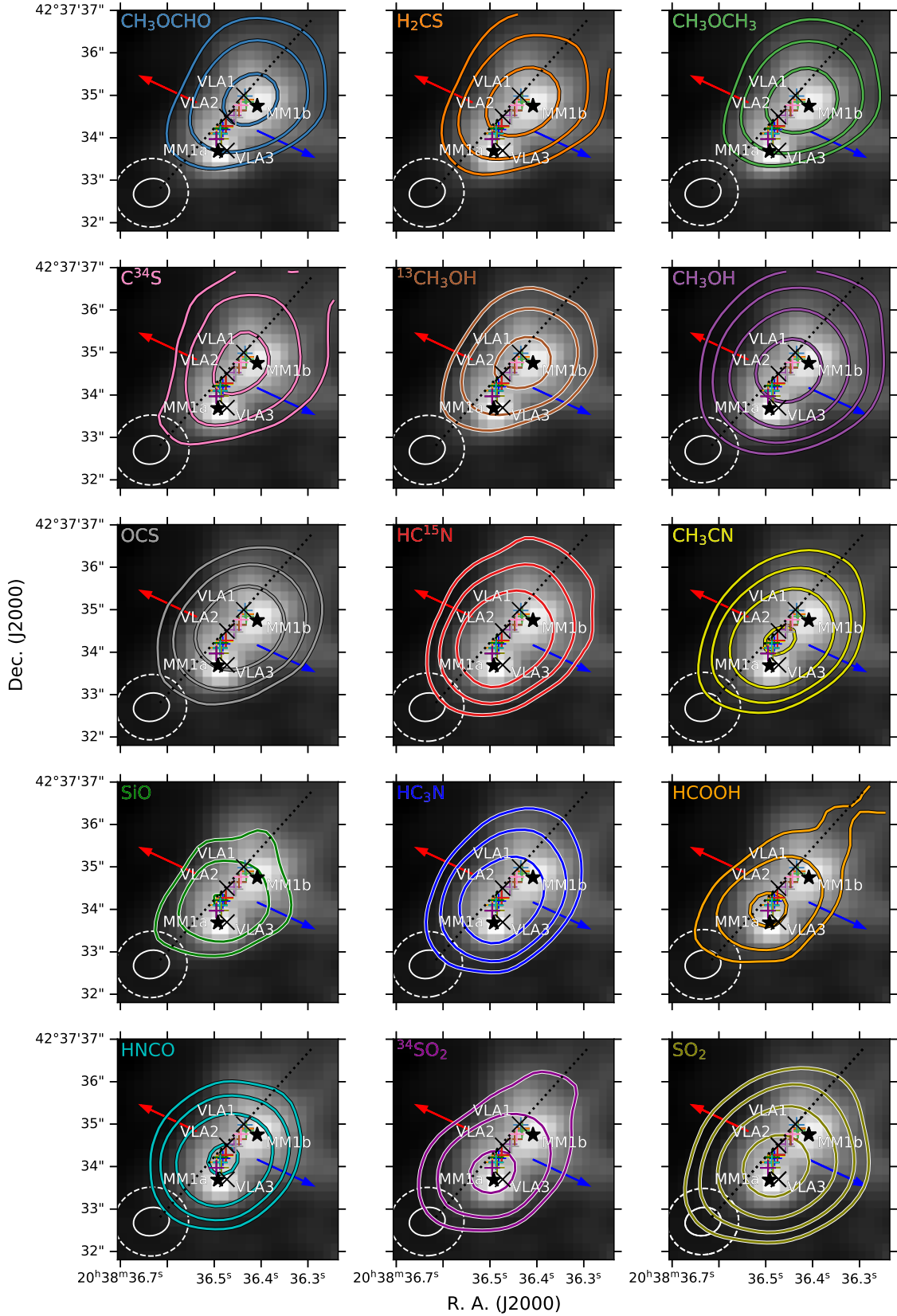


Fig. 11. N30 molecules in contours over continuum in grey-scale. The peak positions of the 2D Gaussian fits to the continuum cores, MM1a and MM1b, are represented with black stars, while the VLA sources are shown with crosses. The molecular peak positions are marked with coloured plus symbols of the same colour as the respective molecule contour plot. The contour levels are 3σ , 6σ , 12σ , and 24σ . The beam sizes are represented by the inner and outer ellipses, with inner representing the continuum, with dimensions as in Fig. 1, and the outer, the beam size of the molecular emission map, which have dimensions of $\sim 1''.65 \times 1''.55$, and position angle of $\sim 11^\circ$ for all molecules. The dotted line represents the red- and blueshifted H_2CO emission, while the arrows represent large-scale CO emission. The panels are sorted according to position of the peak emission, from top to bottom, and left to right.

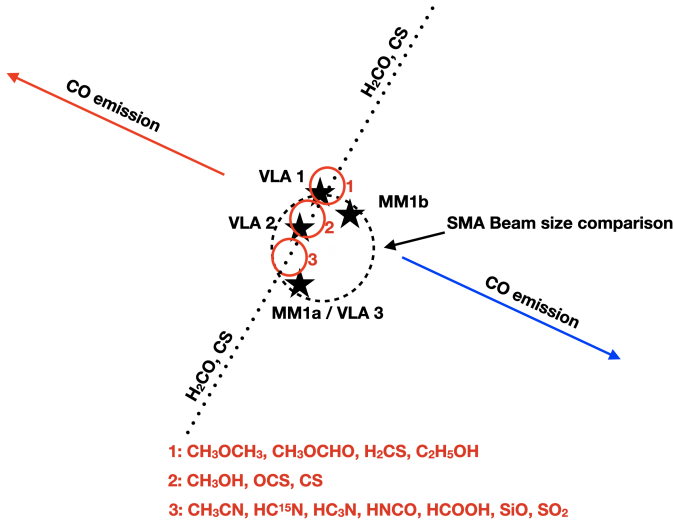


Fig. 12. Representation of the observed molecular gradient towards the MM1 source. The molecules are separated into three groups, with the O-bearing COMs in the first and second groups, with group 1 including H_2CS , and group 2 including CS and OCS . Group 3 includes all the sulphur- and nitrogen-bearing species, as well as SiO and HCOOH .

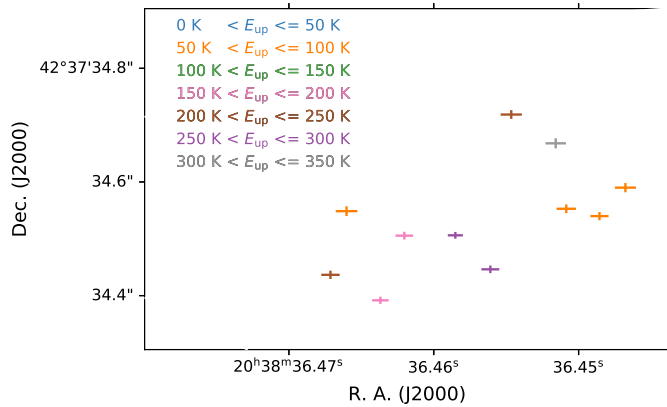


Fig. 13. Peak positions of CH_3OH line transitions, with the upper-level energy of each transition represented in colour. The data points do not follow a systematic pattern.

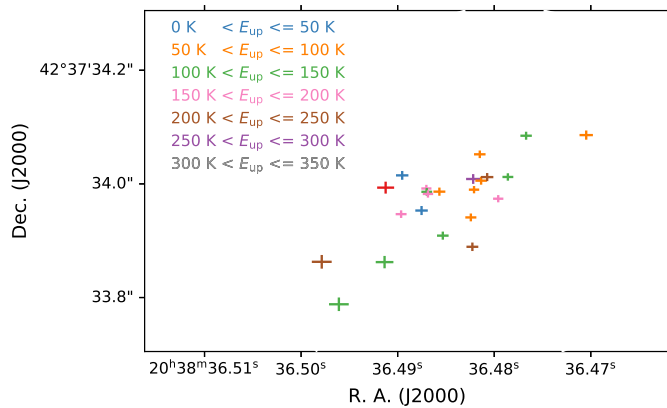


Fig. 14. Peak positions of SO_2 line transitions, with the upper-level energy of each transition represented in colour. As with the CH_3OH line transitions, there does not seem to be any pattern with upper-level energy.

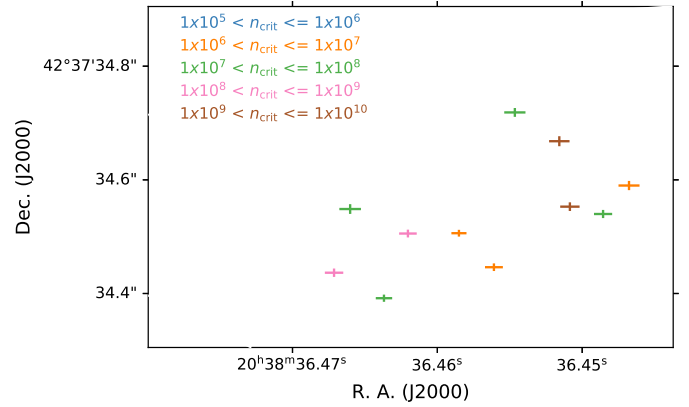


Fig. 15. Peak positions of the CH_3OH line transitions, with the critical density now represented in colour. As with the upper-level energy, there does not seem to be a pattern depending on critical density.

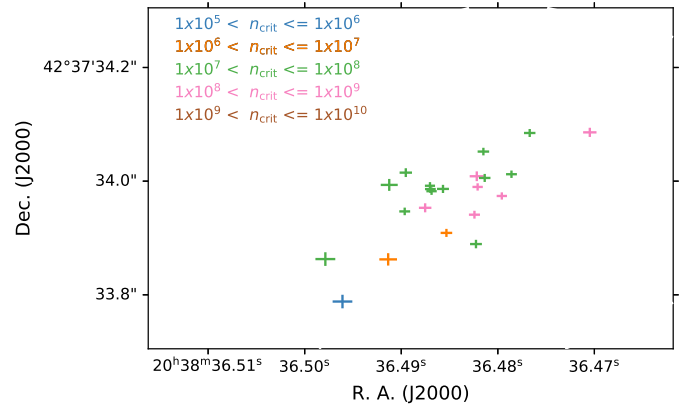


Fig. 16. Peak positions of SO_2 line transitions, with the critical density represented in colour. Again, no systematic pattern is seen.

CH_3CN , we did not detect its isotopologue $^{13}\text{CH}_3\text{CN}$, and so we used the higher energy transition, CH_3CN , $v_8=1$, for the fitting.

Apart from HDO , no other deuterated molecules were detected. Upper limits for the column densities of CH_2DOH were found to be $4.6 \times 10^{16} \text{ cm}^{-2}$, $6.8 \times 10^{16} \text{ cm}^{-2}$, and $8.9 \times 10^{16} \text{ cm}^{-2}$ at positions 1, 2, and 3, respectively. These values were obtained by assuming T_{ex} , v_{source} , and FWHM of $^{13}\text{CH}_3\text{OH}$, and then increasing the column density of the synthetic spectrum until some lines have a strength of $\geq 3\sigma$ of the observed spectrum. The obtained upper limits correspond to a D/H ratio of $\sim 0.1\%$. This value includes statistical correction for the three possible symmetries of the CH_2DOH molecule, which will give an observed $N_{\text{CH}_2\text{DOH}}/N_{\text{CH}_3\text{OH}}$ ratio a factor of three higher than the actual ratio (see e.g. Jørgensen et al. 2018; Manigand et al. 2019).

A number of more complex species have been detected towards high-mass star-forming regions; for example Sgr(B2) near the Galactic centre. One of the more abundant molecules detected towards this source is acetone (CH_3COCH_3) with a relative molecular abundance of X, corresponding to $N(\text{CH}_3\text{COCH}_3)/N(\text{CH}_3\text{OH}) \sim 0.1$ (Belloche et al. 2016). When adopting the same column-density ratio and physical conditions, we find that the non-detection towards N30 is consistent with the noise level.

Relative abundances of the modelled molecules are shown in Figs. 19 and 20, with Fig. 19, showing the column densities

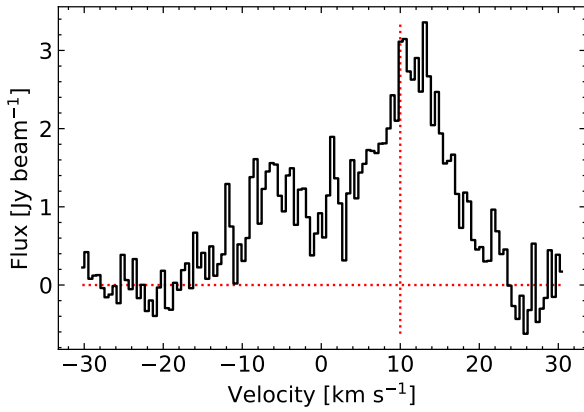


Fig. 17. 347.33 GHz SiO, $J = 8 - 7$ line, showing broad emission, with a weak blueshifted component at -6 km s^{-1} . The horizontal red line represents the baseline, while the vertical line is the source velocity derived for MM1a from the line modelling in CASSIS, at 9.5 km s^{-1} .

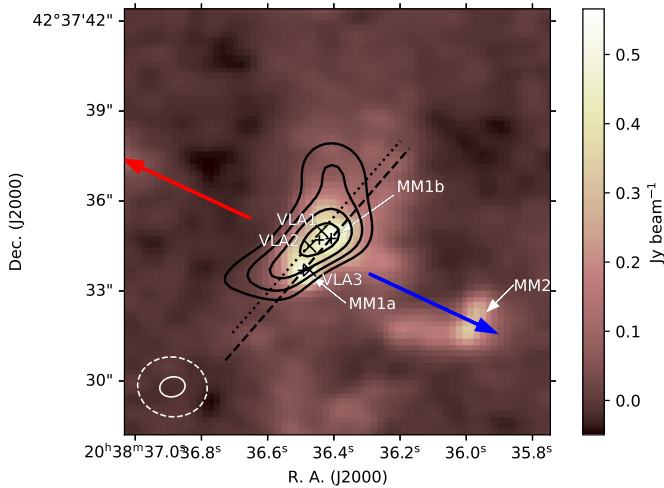


Fig. 18. Continuum image of CygX-N30, with contours showing the 335.395 GHz HDO line emission (transition $J = 3_{3,1} - 4_{2,2}$). The contour levels are at 3σ , 4σ , 5σ and 6σ .

of each molecule with respect to the column density of CH_3OH , at each respective position, whereas Fig. 20 shows the column densities with respect to the column density of CH_3CN . As with the integrated intensity maps, we see how the O-bearing species peak towards position 1, while the N- and S-bearing species peak towards position 3.

4. Discussion

In this section, we discuss our results in more detail, starting with a comparison of the abundances towards N30 with other sources, followed by discussions on excitation temperatures and the D/H ratio observed towards N30. We end the section with a discussion of the continuum cores, and the observed molecular gradient.

4.1. Abundances compared to other sources

To put the inferred abundances into a broader context, we compare them to those observed towards different sources, looking at the differences and similarities. An example of such a

comparison is presented by Drozdovskaya et al. (2019), in which abundances towards the low-mass protostar IRAS 16293B are compared with those measured towards Comet 67P/C-G, with the cometary data used as a representation of the initial ingredients that formed the Earth and the other Solar System planets. From their results, the authors concluded that the volatile composition of comets and planetesimals is partially inherited from the pre- and protostellar phases of the evolution. Jørgensen et al. (2020) extended this comparison to include sources of different masses and located in different environments. They compared, for example, IRAS 16293B and HH212 (both low-mass sources located relatively close to the Sun), AFGL 4176 (a high-mass source also in our Galactic neighbourhood), and Sgr B2 (N2), a high-mass source located close to the Galactic centre. They also included the shocked regions L1157 B1 and Orion KL as well as Comet 67P/C-G in their comparison. The result of this comparison was that the abundances towards these sources are in general very similar, which is an indication that the underlying chemistry is relatively independent of the differences in their physical conditions. There are some differences, however. These latter authors found, for example, that abundances observed towards the protostellar sources tend to agree better with one another than with those towards Comet 67P/C-G, which suggests that some chemical processing occurs between the protostellar and cometary stages (see also Drozdovskaya et al. 2019). In this section, we compare the available column densities observed towards the sources from Jørgensen et al. (2020) to the those we observe towards N30. The column densities taken are relative to CH_3OH , CH_3CN , and SO_2 , representing COM, N-, and S-bearing species, respectively. These sources were selected simply because of the amount of data available in the literature for each of them. While there are many sources in the literature with available column densities, it is rare to find sources observed with wide enough frequency coverage so that column densities can be found for a large amount of molecules. We therefore compare N30 to these sources. As N30 shows a strong chemical gradient, it is important to place the column densities measured towards this source in context with those of the sources from these studies, and in this way achieve a better understanding of the physical nature of the observed chemical gradient.

Figure 21 shows the column density ratios observed towards N30 compared with those observed towards the sources discussed above. The figure shows the column density ratios towards position 2 (see Fig. 12). Similar figures are provided for positions 1 and 3 in Appendix B. The panels in the figure show the column densities of each molecule with respect to CH_3OH , with N30 represented on the y-axis, and each respective source represented on the x-axis. The upper-left panel shows the Galactic centre source, Sgr B2 (N2), which is part of the Sgr B2 molecular cloud, one of the most active high-mass star forming regions in the Galaxy; it was part of the high-resolution EMOCA survey (Exploring Molecular Complexity with ALMA; Bellocche et al. 2016, 2017; Müller et al. 2016; Bonfand et al. 2019). For this source, the O-bearing species show similar abundances (differences within one order of magnitude), while the N-bearing species observed towards N30 have lower ratios compared to CH_3OH (typically by a factor 10).

The upper-right panel shows abundances measured by the ROSINA instrument of the Rosetta mission towards Comet 67P/C-G, as presented by Drozdovskaya et al. (2019) in their comparison to the low-mass source IRAS 16293B which was the target of the PILS program by Jørgensen et al. (2016). Abundances towards Comet 67P/C-G are in general higher than

Table 5. Column densities, excitation temperatures, FWHM and v_{peak} derived at the peak positions of H₂CS and other COMs, excluding CH₃OH (position 1 in Fig. 12).

Molecule	Column density (cm ⁻²)	T_{ex} (K)	$FWHM$ (km s ⁻¹)	v_{peak} (km s ⁻¹)	Peak position ^(a)
COMs and O-bearing species					
CH ₃ OH ^(b)	$1.3(0.2) \times 10^{19}$	140 (25)	5.0	8.5	2
¹³ CH ₃ OH	$1.7(0.2) \times 10^{17}$	140 (25)	5.0	8.5	1/2
CH ₃ ¹⁸ OH ^(c)	$<3.9 \times 10^{16}$	140	5.0	8.5	–
CH ₂ DOH ^(c)	$<5.2 \times 10^{16}$	140	5.0	8.5	–
C ₂ H ₅ OH	$7.2(2.0) \times 10^{16}$	120 (60)	4.5	9.0	1
CH ₃ OCHO	$1.9(0.2) \times 10^{17}$	110 (20)	4.5	8.5	1
CH ₃ OCH ₃	$2.9(0.2) \times 10^{17}$	100 (20)	4.0	8.5	1
H ₂ CO ^(b)	$9.2(1.5) \times 10^{17}$	190 (40)	7.5	8.0	1
H ₂ ¹³ CO	$1.2(0.2) \times 10^{16}$	190 (40)	7.5	8.0	1
t-HCOOH	$2.8(0.5) \times 10^{16}$	190 (50)	7.0	11.0	3
c-HCOOH ^(c)	$<2.3 \times 10^{15}$	190	7.0	11.0	3
HDO ^(d)	$4.6(2.0) \times 10^{16}$	140 (25)	4.0	10.5	–
S-bearing species					
H ₂ CS	$3.3(0.5) \times 10^{16}$	170 (50)	5.5	9.0	1
³³ SO ^(e)	$4.0(2.0) \times 10^{15}$	180 (40)	6.5	10.0	3
SO ₂ ^(b)	$5.9(0.2) \times 10^{17}$	180 (40)	6.5	10.0	3
³³ SO ₂ ^(c)	$<6.1 \times 10^{15}$	180 (40)	6.5	10.0	–
³⁴ SO ₂	$2.7(1.0) \times 10^{16}$	180 (40)	6.5	10.0	3
OCS ^(b)	$3.1(0.4) \times 10^{17}$	170 (50)	5.5	9.5	2
OC ³⁴ S	$1.4(0.2) \times 10^{16}$	170 (50)	5.5	9.5	1
N-bearing species					
HNCO	$4.0(1.0) \times 10^{16}$	120 (20)	6.5	9.5	3
HC ₃ N	$3.0(1.0) \times 10^{15}$	200 (40)	7.0	9.5	3
CH ₃ CN, $v_8=1$ ^(f)	$3.6(2.0) \times 10^{16}$	140 (15)	7.0	10.5	2/3
¹³ CH ₃ CN ^(c)	$<6.0 \times 10^{14}$	140	7.0	10.5	–

Notes. Values are derived from synthetic spectrum fitting with CASSIS, with the assumption of LTE. ^(a)Positions of peak molecular emission are as represented in Fig. 12. ^(b)Column densities for optically thick lines derived using ISM ratios from Wilson & Rood (1994). ^(c)Upper limits were determined by setting T_{ex} , FWHM, and v_{peak} , equal to that of a detected isotopologue. ^(d)To estimate a column density for HDO, the excitation temperature of methanol was assumed. ^(e)Both SO and ³⁴SO had optically thick lines, so only the column density of ³³SO could be derived, using T_{ex} of ³⁴SO₂. ^(f)CH₃CN, $v_8=1$, was used for the model fit of CH₃CN because it is optically thick, and ¹³CH₃CN was not detected.

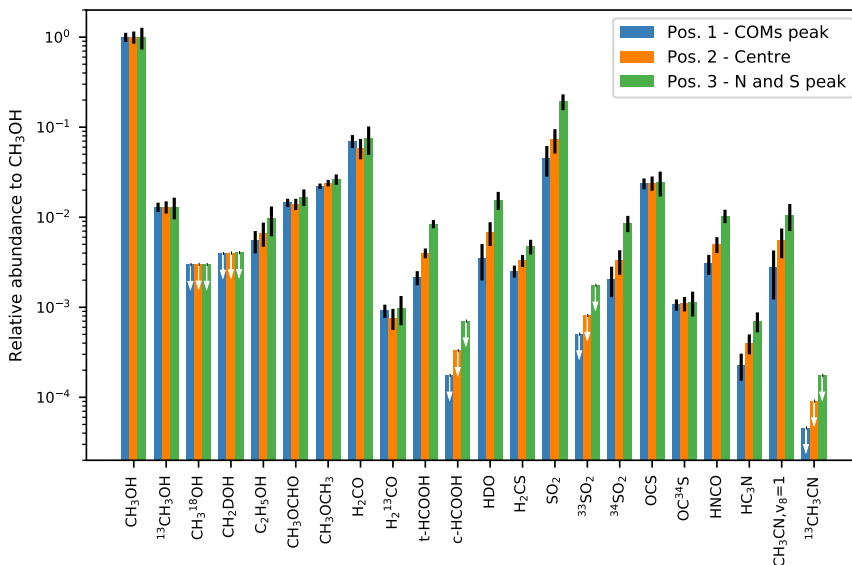
**Fig. 19.** Relative abundances of each molecule relative to the column density of CH₃OH at each respective position. The white arrows represent upper limits, whereas the black bars represent the uncertainties (see Tables 5, 6 and 7).

Table 6. Column densities, excitation temperatures, FWHM and v_{peak} derived at the peak positions of CH₃OH (position 2 in Fig. 12).

Molecule	Column density (cm ⁻²)	T_{ex} (K)	$FWHM$ (km s ⁻¹)	v_{peak} (km s ⁻¹)	Peak position ^(a)
COMs and O-bearing species					
CH ₃ OH ^(b)	$1.0 (0.2) \times 10^{19}$	120 (25)	4.5	9.0	2
¹³ CH ₃ OH	$1.3(0.2) \times 10^{17}$	120 (25)	4.5	9.0	1/2
CH ₃ ¹⁸ OH ^(c)	$<3.0 \times 10^{16}$	100	4.5	9.0	–
CH ₂ DOH ^(c)	$<4.0 \times 10^{16}$	100	4.5	9.0	–
C ₂ H ₅ OH	$6.7(2.0) \times 10^{16}$	130 (60)	5.0	9.5	1
CH ₃ OCHO	$1.4(0.2) \times 10^{17}$	110 (20)	4.5	9.0	1
CH ₃ OCH ₃	$2.4(0.2) \times 10^{17}$	110 (20)	5.0	9.0	1
H ₂ CO ^(b)	$5.9(1.5) \times 10^{17}$	160 (40)	8.0	9.0	1
H ₂ ¹³ CO	$7.6(2.0) \times 10^{15}$	160 (40)	8.0	9.0	1
t-HCOOH	$4.0(0.5) \times 10^{16}$	190 (50)	7.0	11.0	3
c-HCOOH ^(c)	$<3.3 \times 10^{15}$	190	7.0	11.0	–
HDO ^(d)	$6.8(2.0) \times 10^{16}$	120 (25)	4.0	10.5	–
S-bearing species					
H ₂ CS	$3.3(0.5) \times 10^{16}$	140 (50)	5.5	9.0	1
³³ SO ^(e)	$5.4(2.0) \times 10^{15}$	130 (40)	6.5	10.0	3
SO ₂ ^(b)	$7.3(0.2) \times 10^{17}$	130 (40)	6.0	10.0	3
³³ SO ₂ ^(c)	$<8.1 \times 10^{15}$	130 (40)	6.0	10.0	–
³⁴ SO ₂	$3.3(1.0) \times 10^{16}$	130 (40)	6.0	10.0	3
OCS ^(a)	$2.4(0.4) \times 10^{17}$	170 (50)	5.0	10.0	2
OC ³⁴ S	$1.1(0.2) \times 10^{16}$	170 (50)	5.0	10.0	1
N-bearing species					
HNCO	$5.0(1.0) \times 10^{16}$	150 (20)	6.5	10.5	3
HC ₃ N	$4.0(1.0) \times 10^{15}$	210 (40)	7.0	10.0	3
CH ₃ CN, $v_8=1$ ^(f)	$5.5(2.0) \times 10^{16}$	140 (10)	7.0	10.5	2/3
¹³ CH ₃ CN ^(c)	$<9.1 \times 10^{14}$	140	7.0	10.5	–

Notes. Values are derived from synthetic spectrum fitting with CASSIS, with the assumption of LTE. ^(a)Positions of peak molecular emission are as represented in Fig. 12. ^(b)Column densities for optically thick lines derived using ISM ratios from Wilson & Rood (1994). ^(c)Upper limits were determined by setting T_{ex} , $FWHM$, and v_{peak} , equal to that of a detected isotopologue. ^(d)To estimate a column density for HDO, the excitation temperature of methanol was assumed. ^(e)Both SO and ³⁴SO had optically thick lines, so only the column density of ³³SO could be derived, using T_{ex} of ³⁴SO₂. ^(f)CH₃CN, $v_8=1$, was used for the model fit of CH₃CN because it is optically thick, and ¹³CH₃CN was not detected.

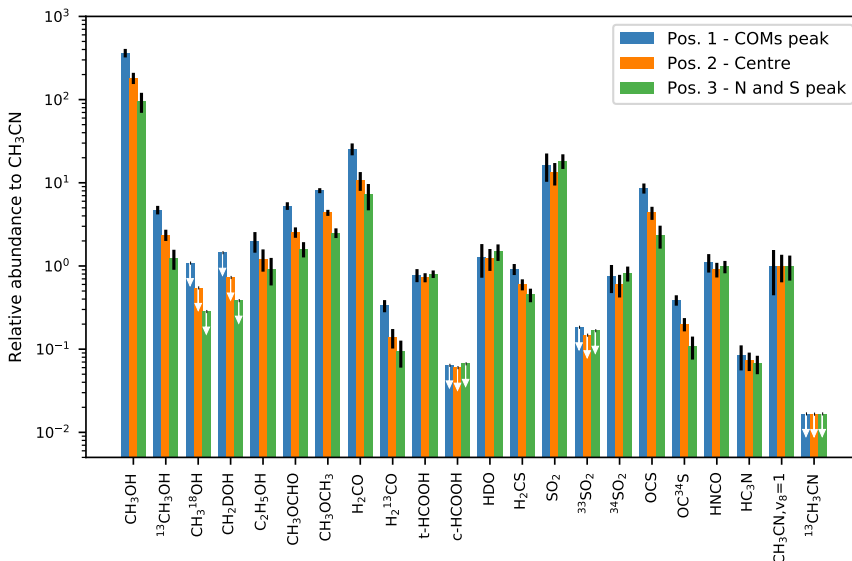
**Fig. 20.** Relative abundances of each molecule relative to the column density of CH₃CN at each respective position. The white arrows and black bars are as in Fig. 19.

Table 7. Column densities, excitation temperatures, FWHM, and v_{peak} derived at the peak positions of the N- and S- bearing species peak (position 3 in Fig. 12).

Molecule	Column density (cm^{-2})	T_{ex} (K)	$FWHM$ (km s^{-1})	v_{peak} (km s^{-1})	Peak position ^(a)
COMs and O-bearing species					
CH_3OH ^(b)	$5.7 (1.5) \times 10^{18}$	100 (25)	5.0	9.5	2
$^{13}\text{CH}_3\text{OH}$	$7.4(2.0) \times 10^{16}$	100 (25)	5.0	9.5	1/2
$\text{CH}_3^{18}\text{OH}$ ^(c)	$<1.7 \times 10^{16}$	100	5.0	9.5	–
CH_2DOH ^(c)	$<2.3 \times 10^{16}$	100	5.0	9.5	–
$\text{C}_2\text{H}_5\text{OH}$	$5.5(2.0) \times 10^{16}$	140 (60)	5.5	10.5	1
CH_3OCHO	$9.6(2.0) \times 10^{16}$	110 (20)	5.0	9.0	1
CH_3OCH_3	$1.5(0.2) \times 10^{17}$	100 (20)	4.5	9.5	1
H_2CO ^(b)	$4.3(1.5) \times 10^{17}$	150 (40)	8.5	9.5	1/2
H_2^{13}CO	$5.6(2.0) \times 10^{15}$	150 (40)	8.5	9.5	1
t-HCOOH	$4.8(0.5) \times 10^{16}$	170 (50)	7.0	11.0	3
c-HCOOH	$<4.0 \times 10^{15}$	170 (50)	7.0	11.0	3
HDO ^(d)	$8.9(2.0) \times 10^{16}$	100 (25)	4.0	11.0	–
S-bearing species					
H_2CS	$2.4(0.5) \times 10^{16}$	140 (50)	5.5	9.5	1
^{33}SO ^(e)	$7.0(2.0) \times 10^{15}$	130 (40)	6.5	10.0	3
SO_2 ^(b)	$1.1(0.2) \times 10^{18}$	130 (40)	6.5	10.0	3
$^{33}\text{SO}_2$ ^(c)	$<1.2 \times 10^{16}$	130	6.5	10.0	–
$^{34}\text{SO}_2$	$4.8(1.0) \times 10^{16}$	130 (40)	6.5	10.0	3
OCS ^(a)	$1.4(0.4) \times 10^{17}$	150 (50)	5.0	10.0	2
OC^{34}S	$6.5(2.0) \times 10^{15}$	170 (50)	5.0	10.0	1
N-bearing species					
HNCO	$5.9(1.0) \times 10^{16}$	160 (40)	6.5	10.5	3
HC_3N	$4.0(1.0) \times 10^{15}$	240 (40)	7.0	10.0	3
CH_3CN , $v_8=1$ ^(f)	$6.0(2.0) \times 10^{16}$	140 (10)	7.0	10.5	2/3
$^{13}\text{CH}_3\text{CN}$ ^(c)	$<1.0 \times 10^{15}$	140	7.0	10.5	–

Notes. Values are derived from synthetic spectrum fitting with CASSIS, with the assumption of LTE. ^(a)Positions of peak molecular emission are as represented in Fig. 12. ^(b)Column densities for optically thick lines derived using ISM ratios from Wilson & Rood (1994). ^(c)Upper limits were determined by setting T_{ex} , $FWHM$, and v_{peak} , equal to that of a detected isotopologue. ^(d)To estimate a column density for HDO, the excitation temperature of methanol was assumed. ^(e)Both SO and ^{34}SO had optically thick lines, so only the column density of ^{33}SO could be derived, using T_{ex} of $^{34}\text{SO}_2$. ^(f) CH_3CN , $v_8=1$, was used for the model fit of CH_3CN because it is optically thick, and $^{13}\text{CH}_3\text{CN}$ was not detected.

that observed towards N30, with no real differences between O- and N-bearing species. However, for IRAS 16293B, the O-bearing species are very similar to N30, while the N- and S-bearing species are in general higher towards N30. This is in contrast to the Galactic centre source Sgr B2 (N2), which also shows similar O-bearing species, but with higher abundances in N-bearing species. This would suggest that N30 has abundances in N-bearing species somewhere in between the Galactic centre (high-mass) source, and the solar neighbourhood (low-mass) source IRAS 16293B.

Orion KL, which is not a ‘traditional’ hot core but is believed to be the result of an explosive event where a pre-existing dense region is heated from the outside (Friedel & Snyder 2008; Zapata et al. 2011; Crockett et al. 2015; Pagani et al. 2017), shows, in general, similar abundances to those observed towards N30. The high-mass source AFGL 4176 (Bøgelund et al. 2019) shows abundances in even better agreement to N30. HH212 is a low-mass source, with COMs observed in its disc atmosphere, that is, above and below the disc (Lee et al. 2019). This source also shows similarities to N30, although the available data in the literature for this source are still limited. L1157-B1 (Arce et al.

2008; Sugimura et al. 2011; Lefloch et al. 2017) on the other hand is a shocked region, with the available column densities for this source suggesting that the O-bearing species are a little less abundant in N30, while the only N-bearing species available for comparison, CH_3CN , is significantly more abundant in N30. Nevertheless, when all the sources are compared, the abundances observed towards AFGL 4176 agree best with those towards N30. At a distance of ~ 3.7 kpc, this source is more distant than N30 ($d \sim 1.3$ kpc), and more luminous, with an upper limit of $2 \times 10^5 L_{\odot}$ (compared to N30, with a luminosity of $\sim 2.04 \times 10^4 L_{\odot}$; Cao et al. 2019). It also has a large disc-like structure, as was found by Johnston et al. (2015) using ALMA observations of CH_3CN on scales of ~ 1200 AU. Furthermore, Bøgelund et al. (2019) found (also from ALMA observations, at a resolution of ~ 1285 AU) that the O- and N-bearing species in this source also peak in slightly different locations, with N-bearing species peaking slightly closer to the continuum peak than the O-bearing species. This would suggest that a combination of processes could be causing the sublimation of the COMs from the dust grains rather than the traditional hot core model. The agreement with Orion KL also strengthens the case that

it is a combination of processes that result in the COMs we observe.

It therefore seems that, when comparing the abundances observed towards N30, they agree more with other high-mass sources, such as AFGL 4176 (a more traditional hot core, but with some chemical differentiation) and Orion KL (a high-mass, shocked region), than with the abundances observed towards low-mass sources, such as IRAS 16293B, or the outflow region, L1157-B1. The agreement is also better with these high-mass sources than compared to high-mass sources in the Galactic centre. The environment here, which has a cosmic ray ionisation rate of a factor of 50 higher than usually assumed for the Galactic disc (Guzmán et al. 2015; Bonfand et al. 2019), resulting in a higher dust temperature. This presents some challenges for our understanding of the formation of COMs (Jørgensen et al. 2020), which are believed to form primarily on grain surfaces at low temperatures. However, it might explain the high abundance of N-bearing species in the Galactic centre, which evolves over longer timescales in the gas phase (Garrod et al. 2017; Bøgelund et al. 2019). It might also explain the low abundance of S- and N-bearing species in low-mass sources, where the temperatures are lower than in their high-mass counterparts. Bøgelund et al. (2019) suggest that AFGL 4176 is a very young source, where little processing of the chemical inventory by the protostar has occurred, which is why they observe N-bearing species to be low in abundance compared to Sgr B2. For the shocked region L1157-B1, the O-bearing species are sputtered off of the dust grains, but there is not sufficient time for the nitrogen chemistry to evolve in the gas phase (see chemical models by e.g. Garrod 2013; Barone et al. 2015; Garrod et al. 2017; Codella et al. 2017). The agreement between the abundances towards N30 MM1 and AFGL 4176 would suggest that the sources are at a similar stage of evolution, although at scales of ~ 1200 AU, AFGL 4176 appears to be a single source, as opposed to N30, which is at least a binary source at scales of ~ 1300 AU, with the MM1b/VLA1 core probably less evolved than the MM1a/VLA3 core.

Figure 22 shows the abundances of O-bearing species normalised to CH_3OH , with N30 on the x -axis and the respective sources depicted with different symbols on the y -axis. Figure 23 shows the N and S-bearing species normalised to CH_3CN and SO_2 , respectively. Some trends that can be seen in Fig. 22 are that the abundances of O-bearing species in Comet 67P and L1157-B1 are in general higher than the protostellar sources (around one order of magnitude), while the shocked region Orion KL shows higher relative abundances in CH_3OCHO and CH_3OCH_3 (5–10 times higher), but with H_2CO and $\text{C}_2\text{H}_5\text{OH}$ comparable to N30. IRAS 16293B has abundances in O-bearing species that are a little higher than N30 (around five times), while Sgr B2 (N2) has a higher abundance in $\text{C}_2\text{H}_5\text{OH}$ and (five to ten times), and with abundances of CH_3OCHO , CH_3OCH_3 , and H_2CO similar to those for N30. As discussed above for Fig. 21, we also see from this figure that the best agreement is with the high-mass source AFGL 4176. This can also be seen in the abundances with respect to CH_3CN and SO_2 in Fig. 23, with AFGL 4176 showing similar abundances, while Orion KL has higher abundances in N-bearing species but a similar abundance of H_2CS . However, for the N- and S-bearing species, the number of molecular species available for comparison is limited.

Therefore, while N30 clearly shows a strong chemical gradient with different species peaking in different locations, the inferred abundances are typical of what is seen towards other intermediate- and high-mass protostars. This suggests that, no matter the physical origin of the chemical gradient,

the chemistry does not change significantly between different sources.

4.2. Excitation temperatures

We find some differences when comparing the excitation temperatures of different molecular species. The N- and S-bearing species seem to be slightly warmer than the O-bearing species, with ranges from 130 to 200 K and 100 to 140 K, respectively. HC_3N seems to be warmer than other molecular species, with a range of 200–240 K between the three positions in Fig. 12. t-HCOOH also seems to be warmer (170–190 K) than other O-bearing species, but considering that it peaks at the position with the N-bearing species, this might not be strange, because it does not seem to be connected with the other O-bearing molecules. H_2^{13}CO also has a warmer excitation temperature, in the range of 150–190 K.

Other authors have found more significant differences in the excitation temperatures of N- and O-bearing species towards other sources. An example is Orion KL, in which the O-bearing species trace gas of lower temperatures – ranging between 100 and 150 K – than the N-bearing species, which trace gas of 200–300 K (Bell et al. 2014; Crockett et al. 2015). Jørgensen et al. (2018) also found a range of 80–300 K in excitation temperatures of different molecular species in IRAS 16293, and suggested that the differences observed could be attributed to binding energy. van 't Hoff et al. (2020) argues that the so-called ‘soot line’ located close to the protostar at a temperature of ~ 300 K is marked by an excess of hydrocarbons and nitriles sublimated from carbon grains inside this line. The O-bearing COMs on the other hand desorb around the water snowline, which is located further out from the protostar at a temperature of ~ 100 K. However, in the case of N30, we do not observe the COMs concentrated at a central position, but rather spread out in a chemical gradient, which suggest that it is probably not a traditional hot core(s) but rather a combination of factors causing the chemical gradient and the difference in excitation temperature that we observe, including two or more protostars (see Sect. 4.4).

4.3. D/H ratio

We find an upper limit on the D/H ratio derived for $N(\text{CH}_2\text{DOH})/N(\text{CH}_3\text{OH})$ of about 0.1%. Compared to IRAS 16293B, which has values of 2%–3%, this is very low. It is also lower than values reported by Neill et al. (2013) for Orion KL, in which they found similar D/H ratios for water, formaldehyde, and methanol, 0.2%–0.8%. However, for Sgr B2 (N2), Belloche et al. (2016) reported a XD/XH ratio for CH_2DOH of 0.12%, which they argue might be because of the higher kinetic temperatures in the Galactic centre region compared to nearby star-forming regions. Bøgelund et al. (2019) do not detect any deuterated species towards AFGL 4176, which further strengthens the agreement between this source and N30, as discussed in the abundance comparison above. Other high-mass sources have also shown low D/H ratios (e.g. Ratajczak et al. 2011; Bøgelund et al. 2018). Bøgelund et al. (2018) conclude that warm formation temperatures of ~ 30 K could account for the low deuteration in the NGC 6334I regions, with the pre-stellar cloud heated by a nearby O-type star and associated HII region. In the case of N30, the presence of the strong radiation field caused by the nearby Cyg–OB2 association (projected distance of ~ 23 pc at a distance of 1.3 kpc; Reipurth & Schneider 2008) may have heated the pre-stellar cloud and resulted in the low levels of deuteration we observe.

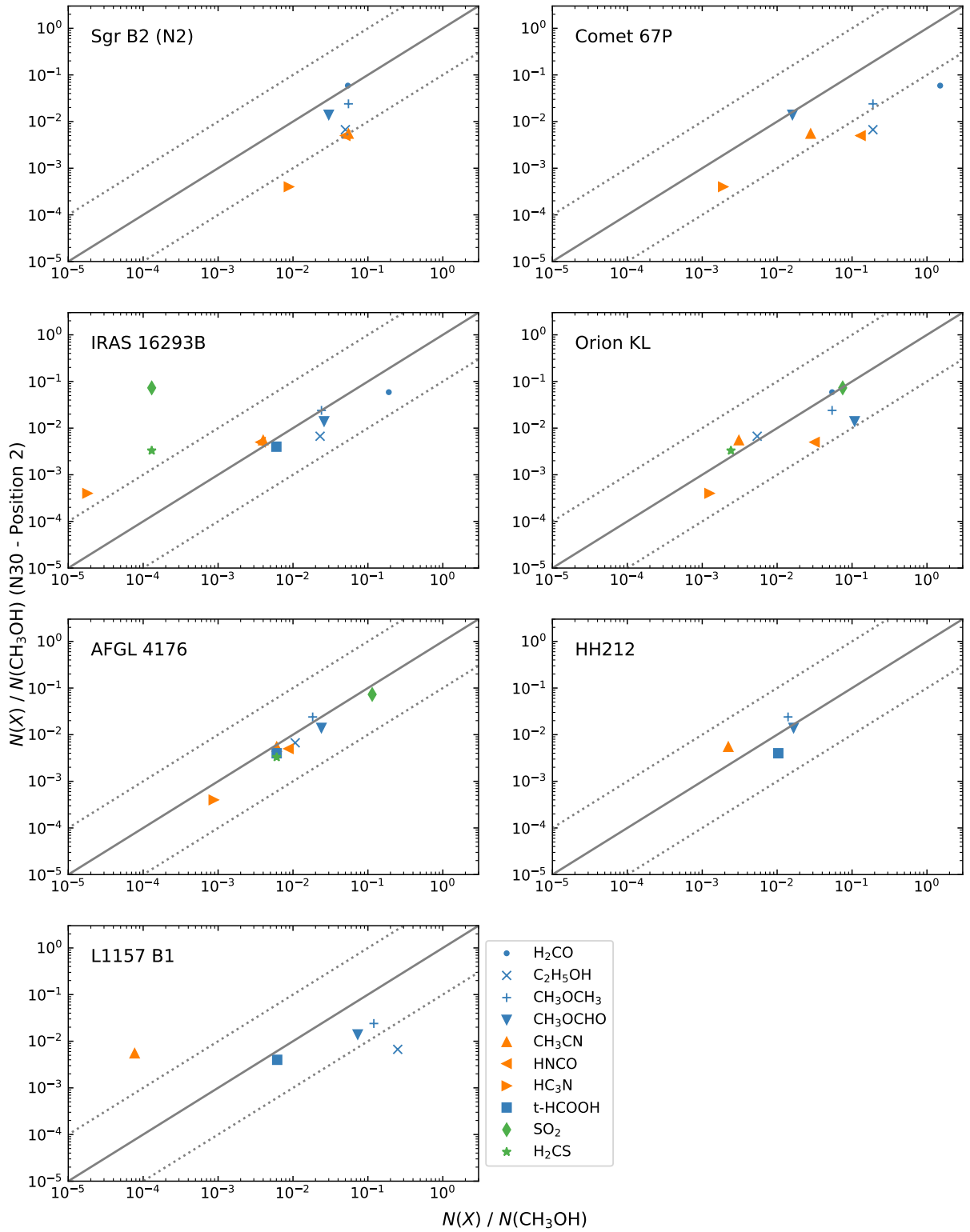


Fig. 21. Column densities towards N30, position 2, with respect to CH_3OH on the y -axis compared to different sources plotted on the x -axis. The solid line represents equal abundances, whereas the dotted lines represent an order of magnitude difference.

An alternative is that the prestellar phase is cold but short, as is expected for higher mass sources. In this scenario, time is the bottleneck in setting the deuteration, as opposed to above where temperature is the limiting factor. Either way, the low degree of deuteration in sources such as this one is not unexpected.

4.4. Continuum cores, and the observed molecular gradient

We find that the O-bearing species have their peak emission concentrated towards the radio (cm) continuum source, VLA1, close to but to the west of the MM1b submm continuum core. S- and N-bearing species on the other hand peak between VLA2 and

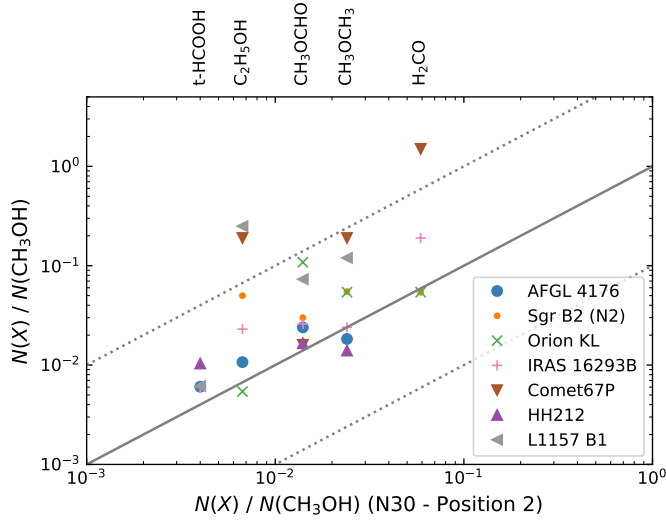


Fig. 22. Column densities with respect to CH_3OH towards N30, position 2, on the x-axis compared to different sources plotted on the y-axis. Each source is marked with a different marker. The solid and dotted lines are as in Fig. 21.

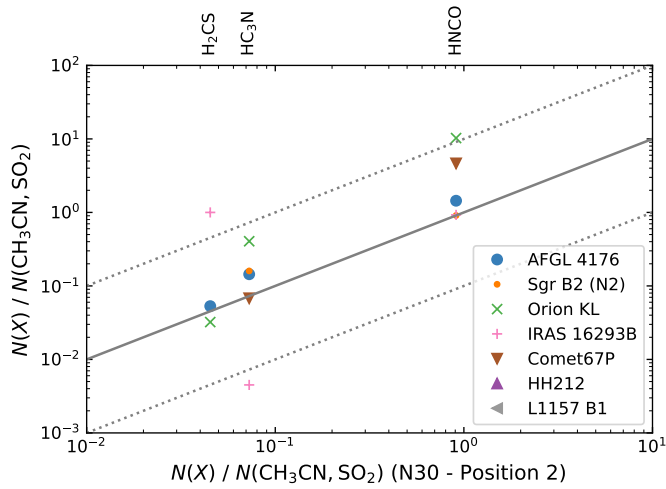


Fig. 23. Column densities towards N30, position 2 (on the x-axis), with respect to CH_3CN for the N-bearing species and SO_2 for H_2CS , compared to different sources plotted on the y-axis. Each source is marked with a different marker. The solid and dotted lines are as in Fig. 21.

VLA3 to the north of MM1a. All molecules peak on a gradient traced by the red- and blueshifted axis of H_2CO and CS emission. This axis is nearly perpendicular to the larger scale (~ 1 pc; Fischer et al. 1985) bipolar outflow axis seen in CO emission, which might indicate that the observed submm and radio continuum sources are located in a disc-like structure (see Hutawarakorn et al. 2002). It is important to note here that this disc-like structure is more akin to a pseudo-disc that is not rotation-supported, but just a rotating structure around the protostars. Looking at the H_2CO and CS maps in Figs. 3 and 6, it is also clear that this structure is very large, with a radius of around 3000 AU, and encompass all the continuum sources (see also Hutawarakorn et al. 2002). Similar disc-like structures are also observed in other sources, such as G35.20-0.74N, in which Hutawarakorn & Cohen (1999) observed a large-scale outflow structure seen in CO emission perpendicular to a disc-like structure containing four hot cores. Chemical differentiation between

the different cores was also observed for this source (with separation between the cores of 1000–2000 AU; Allen et al. 2017), with O- and S-bearing species detected towards the inner two cores (B1 and B2), but few N-bearing species, while the outer two cores (B3 and A) have strong detections of N-, S-, and O-bearing species. Another source is G328.2551–0.5321, for which Csengeri et al. (2018) found N-bearing COMs peaking towards the protostar, while the O-bearing COMs peak towards two spots, offset from the protostar (see also Csengeri et al. 2019). The authors interpreted the two spots as accretion shocks onto a disc with an estimated distance of 200–800 AU from the central protostar. These large disc-like structures are therefore not uncommon in high-mass protostars, and chemical differentiation also seems to be a common feature. Furthermore, with the high angular resolution provided by interferometric observations, more and more examples of sources are identified in which the observed COM emission cannot be explained by the canonical hot-core scenario in which COMs sublimate from the heated dust grains close to the protostar. This was also seen by Bellocche et al. (2020) for low-mass protostars in the CALYPSO survey.

For N30, we observe chemical differentiation between S- and N-bearing species associated with the VLA2 and VLA3/MM1a cores on the one hand, and COMs and O-bearing species associated with the VLA1/MM1b core on the other. Molecules that do not follow this differentiation in O- and S-/N-bearing species are t-HCOOH, observed to peak towards position 3 at the N- and S-bearing species position, and CS, OCS, and H_2CS that peak towards the O-bearing species position.

Our results therefore suggest that N30 is not a purely traditional hot core in which ice-covered dust grains collapse towards a warm protostar and are heated to temperatures ≥ 100 K as they get closer, where the ices sublimate and we see a peak of COM emission (see e.g. Herbst & van Dishoeck 2009, and references therein). The release of the COMs into the gas phase may then lead to further reactions in the dense warm gas, but this is still centred on the region where the dust temperature exceeds ~ 100 K. Instead, the O-bearing COM emission is likely caused by a combination of processes, including accretion of infalling material onto the disc surface – which is similar to what was found by Csengeri et al. (2018) for G328.2551–0.5321 –, while the N- and S-bearing species towards VLA3/MM1a might be a slightly more evolved source where the gas-phase chemistry had more time to evolve.

5. Summary and conclusions

We observed CygX-N30 and MM1 with the SMA in the 345 GHz frequency window at a resolution of $\sim 1''$ and with 32 GHz of continuous frequency coverage. About 400 lines were detected from 29 different molecular species and their isotopologues.

We observe a chemical gradient of molecules along the axis of a disc-like structure perpendicular to the large-scale bi-polar CO emission observed in previous studies. This disc is parallel to but offset from the axis connecting the mm continuum cores (by $\sim 1''$). The O-bearing molecular emission peaks are close to but offset to the west of the MM1b continuum core and are between the VLA1 and VLA2 radio continuum cores, which fall on the molecular gradient-disc axis. The N- and S-bearing species on the other hand are concentrated closer to the MM1a core, between the VLA2 and VLA3 radio continuum cores. The disc-like structure is observed from the red- blueshifted H_2CO and CS emission, which signifies infalling gas onto the surface of the disc-like structure. This implies that the COMs observed

towards MM1 are not purely the result of a traditional hot core, but are rather caused by a combination of processes, including accretion. In order to test this hypothesis, it will be useful to re-observe this source in a few years to check for time variability.

Comparing the abundances observed towards N30 MM1 with other sources, we find that it is similar to the high-mass protostar AFGL 4176, and to a lesser extent to the shocked region Orion KL, and to the Galactic centre source Sgr B2 (N2). The O-bearing species show similar abundances with respect to CH₃OH towards these sources, whereas for Sgr B2 (N2), the N-bearing species are less abundant in N30, while the low-mass source IRAS 16293B has lower abundances in N- and S-bearing species; the O-bearing species are similar. The agreement with AFGL 4176 seems to suggest that N30 MM1 is at a similar stage of evolution to this source, but with the MM1b/VLA1/VLA2 source at the centre of the disc-like structure probably at an earlier stage than the MM1a/VLA3 core, where the N- and S-bearing species had more time to evolve.

We observe a small difference in excitation temperature between the O-bearing species and the S- and N-bearing species. A larger difference is observed for HC₃N, H₂¹³CO, and t-HCOOH, which seem to trace warmer gas than the other species. The low levels of deuteration in N30 MM1 suggest that the grain-surface formation temperature of COMs in the pre-stellar cloud was warm, that is, ~30 K, probably the result of radiation from the nearby Cyg-OB2 association.

In conclusion, these observations highlight the benefit of being able to observe large frequency ranges simultaneously and at high angular resolution in order to shed light on the physical origin of COMs. Particularly, the frequency coverage ensures that a large number of species from different chemical families are covered simultaneously, and the high angular resolution allows spatial correlations between species to be closely examined. This is especially important for the physically complex structure of sources such as N30, which complicates our understanding of the origin of COMs. It is therefore important to study such sources with multiple chemical tracers, and at sufficient angular resolution to be able to disentangle the various physical and chemical processes taking place. Further analysis of the remaining nine sources in the PILS-Cygnus survey will clearly continue to shed light on their origin.

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Appendix A: Observing log

The full observing log is provided in Table A.1. Specifically, this includes the observing dates and calibrators used for each night of the observations. The observing strategy was to observe two science targets between a pair of gain observations, and to loop over all ten targets in each track. This ensured uniform sensitivity. Furthermore, a random source was chosen to start each track, to ensure near uniform uv coverage for all sources, and thus similar image fidelity.

Appendix B: Column density comparison

The column density comparisons with other sources at positions 1 and 3 (see Fig. 12) are provided here, with the comparison at position 2 shown and discussed in the text. There is a slight difference, as expected from the difference in column densities of the different positions, with a slightly better agreement with Orion KL and AFGL 4176 at position 3.

Appendix C: Position–velocity maps

Position–velocity maps of selected molecular lines are presented in Fig. C.1, with the maps of H_2CO and CS shown (Figs. 9 and 10 respectively) and discussed in the text. A full kinematics study of the source is beyond the scope of this work, but it is clear from Fig. C.1 that different molecules trace different regions in the system, which is also seen in the molecular gradient that we observe.

Appendix D: Molecular line plots

In this Appendix we present plots of all the lines for each of the molecules listed in Table 6, taken at position 2 in Fig. 12, for which we made a model fit. In all the plots, the model is overplotted in red over the observed spectrum in black. Only lines detected above $3\sigma \sim 10$ K intensity are shown.

Table A.1. Observing log.

Observing date	No of antennas	Configuration	Bandpass	Flux	Gain	$\tau(225\text{ GHz})$
21/06/2017	7	COM	3c454.3	Titan, Neptune	mwc349a	0.05–0.07
22/06/2017	7	COM	3c273, 3c454.3	Titan, Neptune	mwc349a	0.10
27/06/2017	7	COM	3c454.3	Callisto	mwc349a	0.08
10/07/2017	7	COM	3c454.3	Callisto	mwc349a	0.05–0.06
07/08/2017	6	COM	3c84	Titan, Uranus	mwc349a	0.05
20/10/2017	8	EXT	3c84	Uranus	mwc349a	0.02–0.03
22/10/2017	7	EXT	3c84	Uranus	mwc349a	0.08–0.10
08/11/2017	8	EXT	3c84	Uranus	mwc349a	0.07
09/11/2017	7	EXT	3c84	Uranus	mwc349a	0.07
10/11/2017	8	EXT	3c84	Uranus	mwc349a	0.07

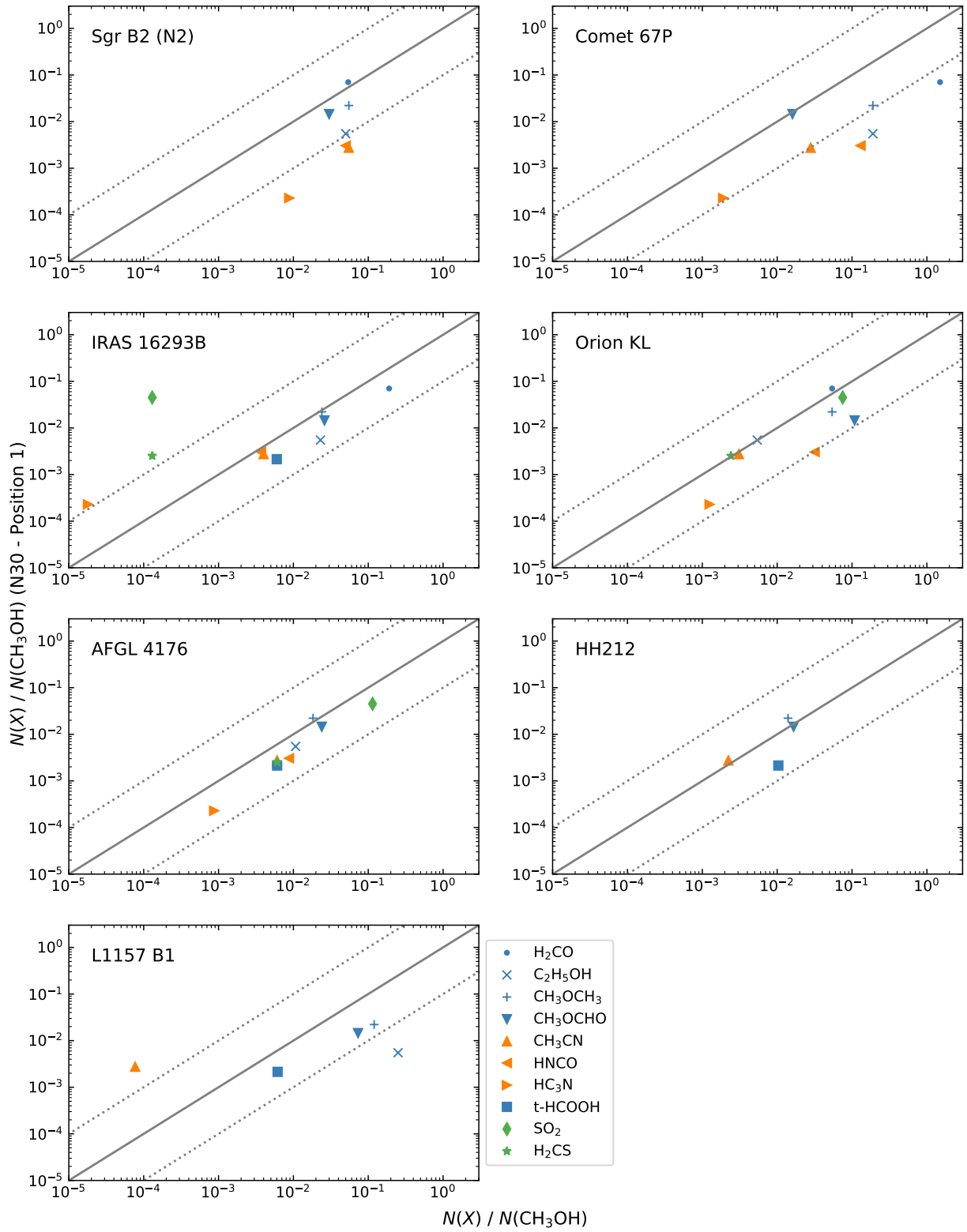


Fig. B.1. Column densities towards N30, position 1, with respect to CH_3OH , on the y-axis, compared to different sources plotted on the x-axis. The solid line represents equal abundances, whereas the dotted lines represent an order of magnitude difference.

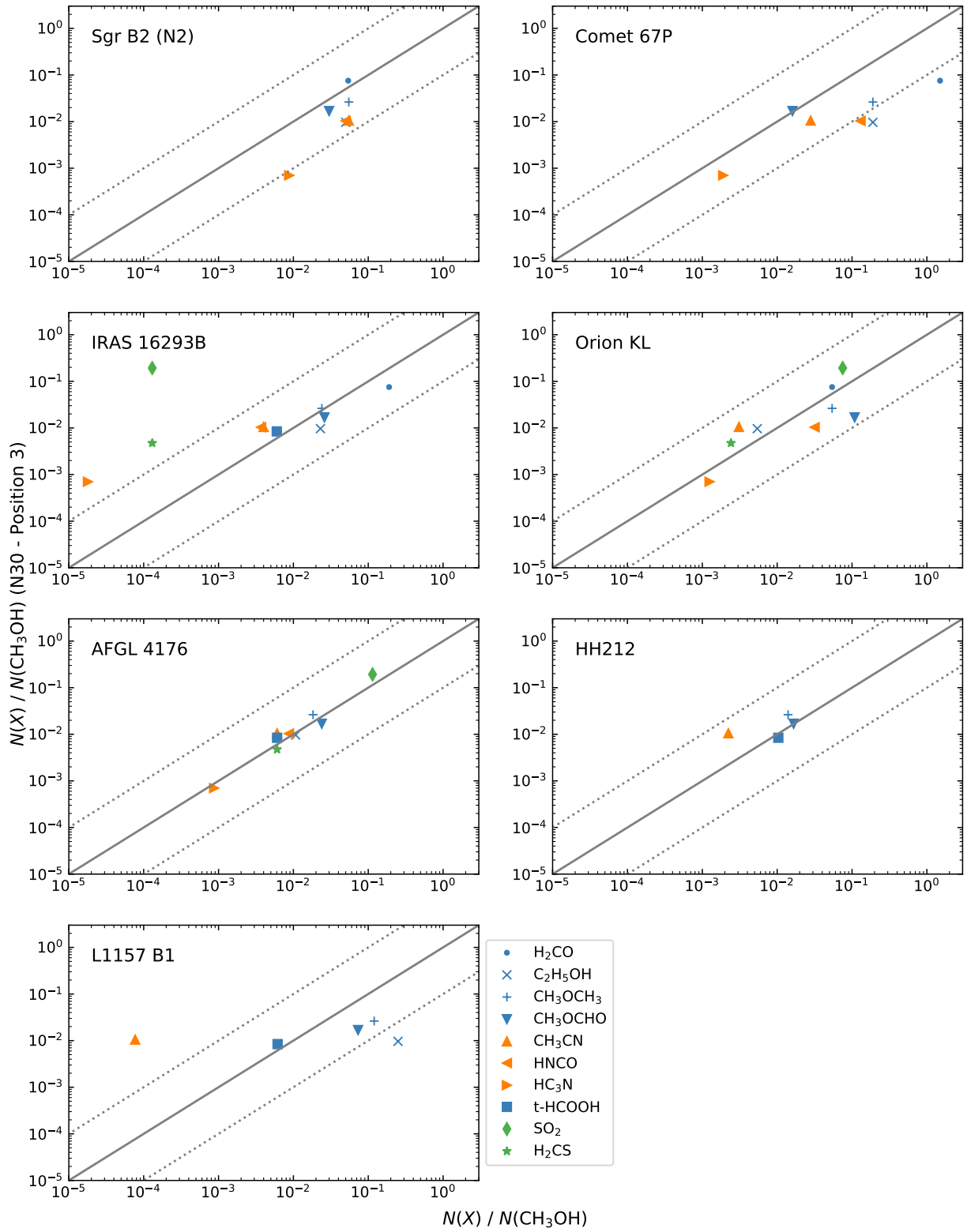


Fig. B.2. Column densities towards N30, position 3, with respect to CH_3OH , on the y-axis, compared to different sources plotted on the x-axis. The solid line represents equal abundances, whereas the dotted lines represent an order of magnitude difference.

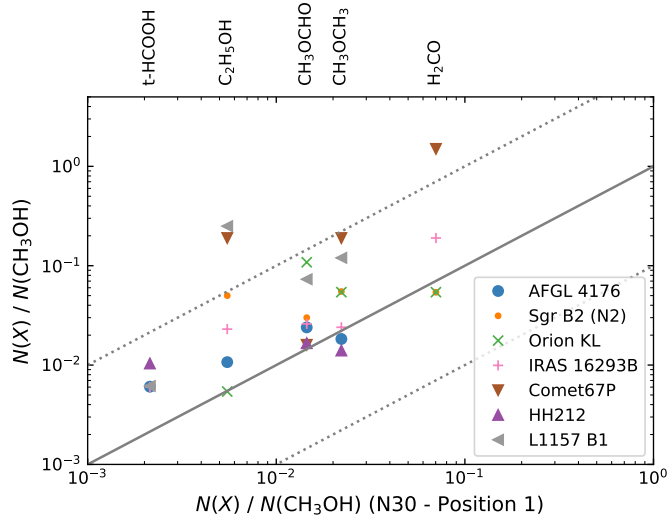


Fig. B.3. Column densities with respect to CH_3OH towards N30, position 1, on the x-axis, compared to different sources plotted on the y-axis. Each source is marked with a different marker. The solid and dotted lines are as in Fig. B.1

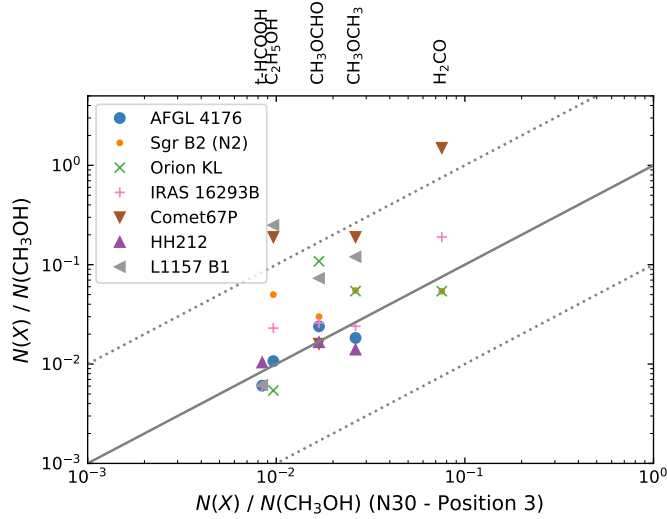


Fig. B.4. Column densities with respect to CH_3OH towards N30, position 3, on the x-axis, compared to different sources plotted on the y-axis. Each source is marked with a different marker. The solid and dotted lines are as in Fig. B.1

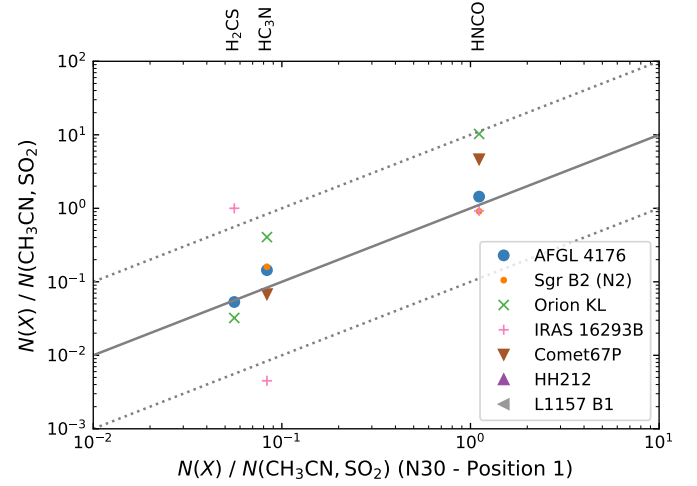


Fig. B.5. Column densities towards N30 position 1 (on the x-axis), with respect to CH_3CN , for the N-bearing species, and SO_2 for H_2CS , compared to different sources plotted on the y-axis. Each source is marked with a different marker. The solid and dotted lines are as in Fig. B.1

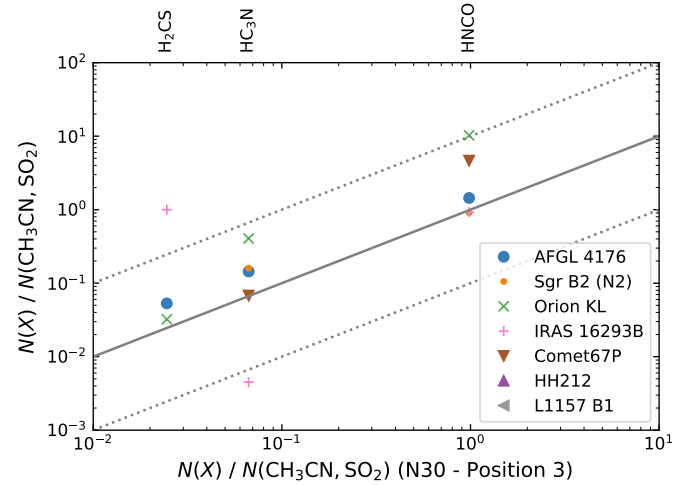


Fig. B.6. Column densities towards N30 position 3 (on the x-axis), with respect to CH_3CN , for the N-bearing species, and SO_2 for H_2CS , compared to different sources plotted on the y-axis. Each source is marked with a different marker. The solid and dotted lines are as in Fig. B.1

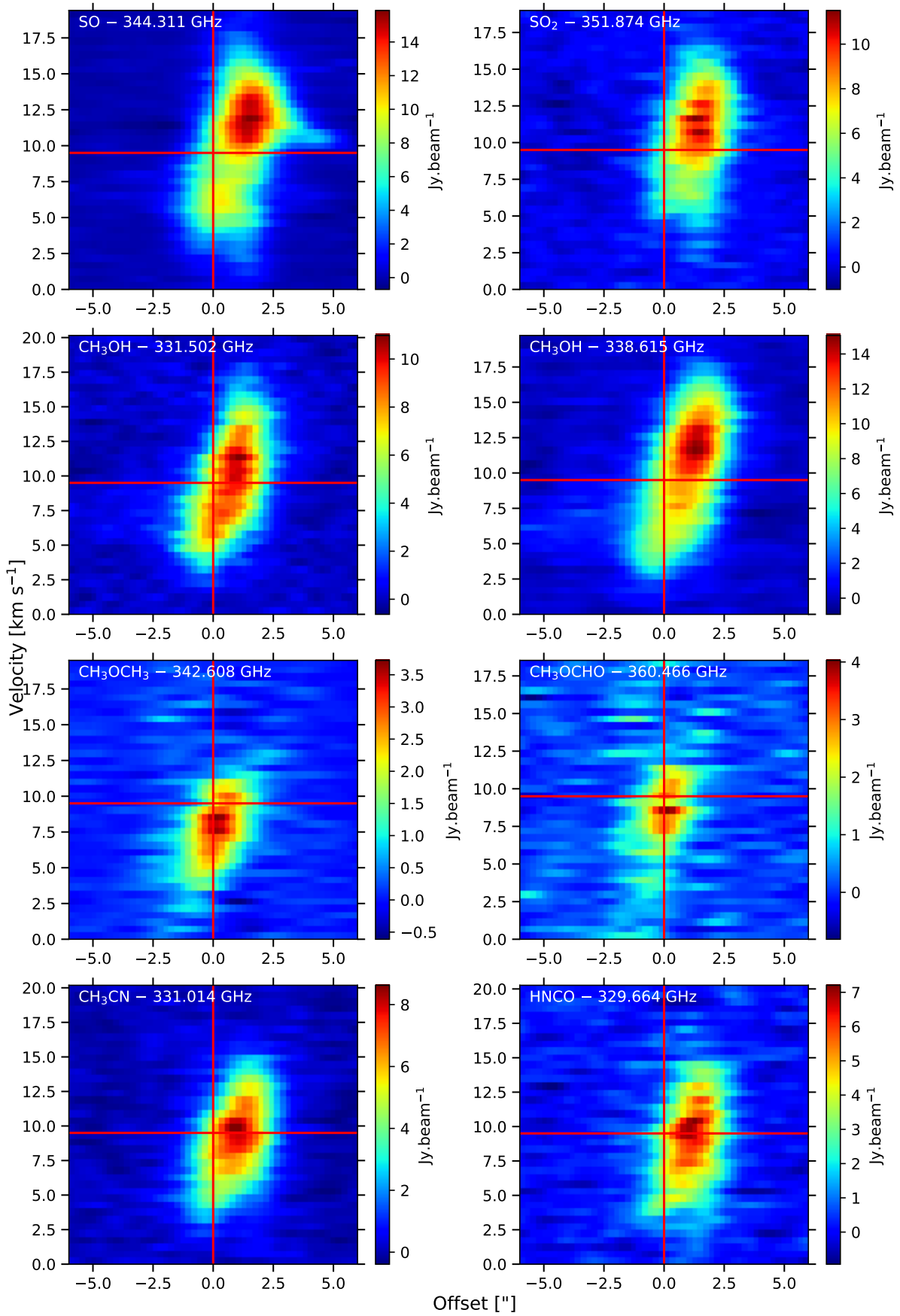


Fig. C.1. Position–velocity maps of selected molecular lines. The red vertical line represents the position of 0'' offset from the centre position between the peak red- and blueshifted H₂CO emission (see 9). The horizontal line represents the systemic velocity, also for H₂CO.

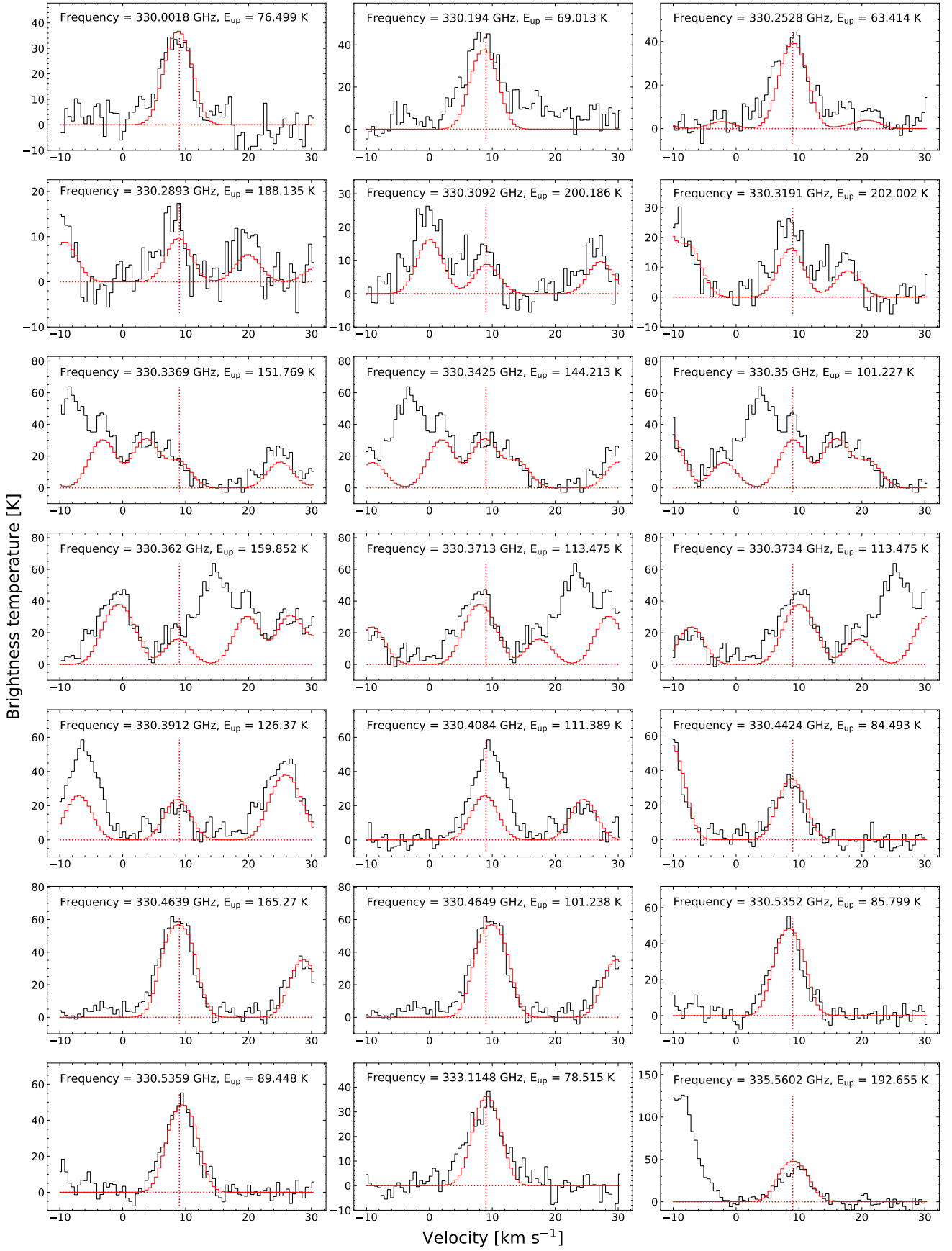


Fig. D.1. $^{13}\text{CH}_3\text{OH}$ synthetic spectrum overplotted on the observed spectrum. The fitted values obtained were column density = $1.3 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 120.0 K, line width (FWHM) = 4.5 km s $^{-1}$, and source velocity (v_{source}) = 9.0 km s $^{-1}$.

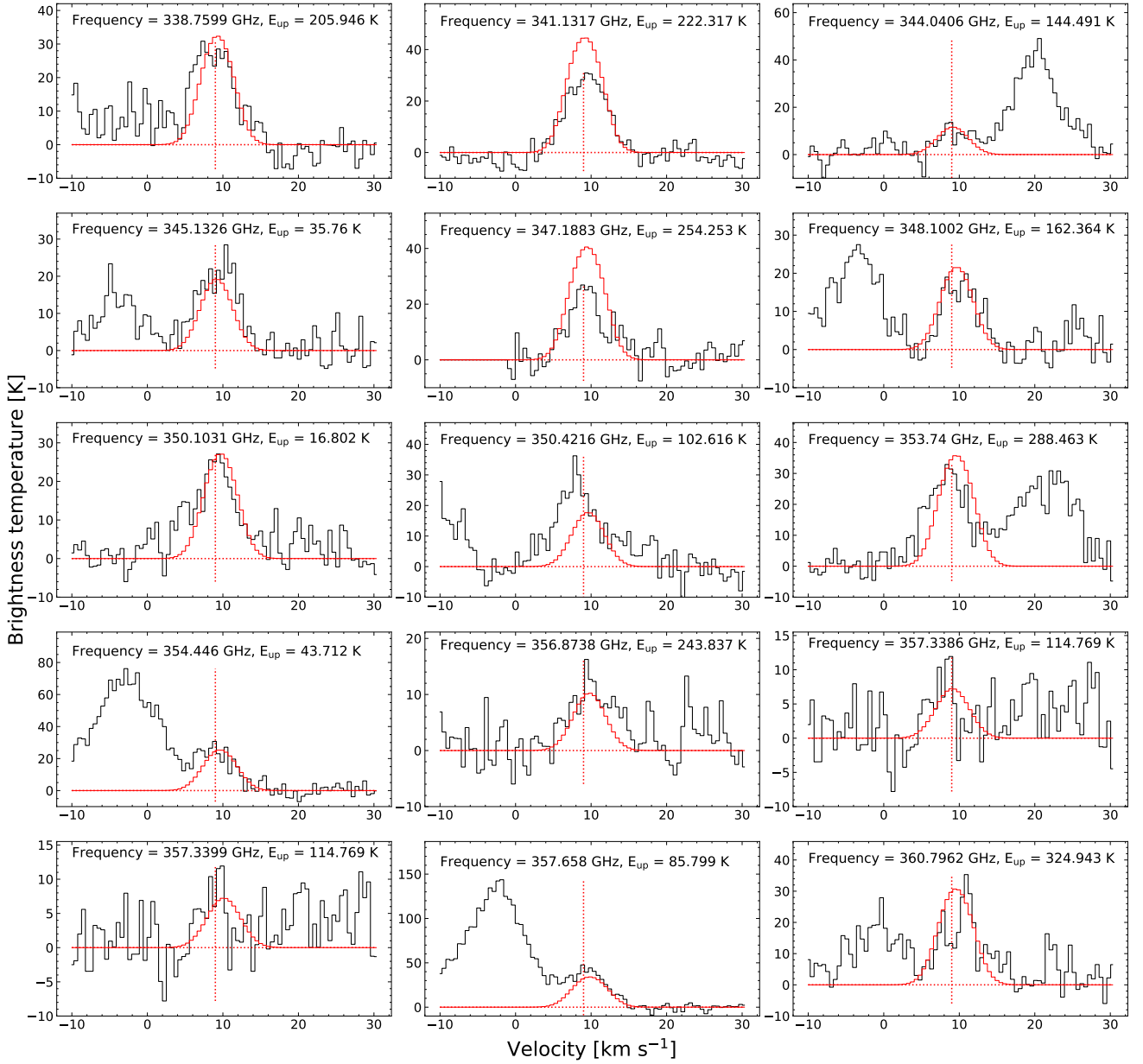


Fig. D.2. $^{13}\text{CH}_3\text{OH}$ synthetic spectrum overplotted on the observed spectrum. The fitted values obtained were column density = $1.3 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 120.0 K, line width (FWHM) = 4.5 km s $^{-1}$, and source velocity (v_{source}) = 9.00 km s $^{-1}$.

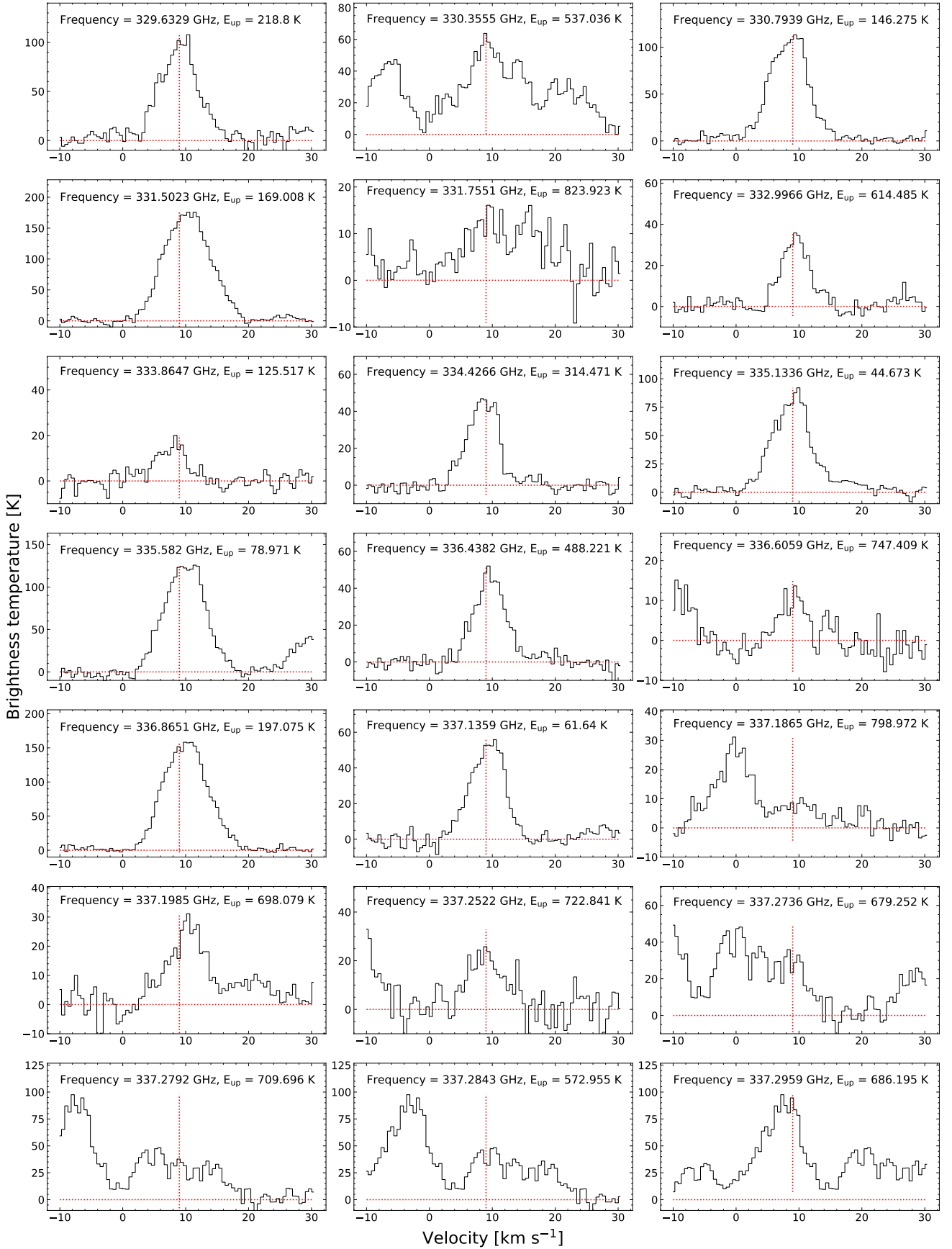


Fig. D.3. CH_3OH observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of $^{13}\text{CH}_3\text{OH}$ with the ISM isotopologue ratio of $^{12}\text{C}/^{13}\text{C} = 77$. The obtained column density was $1.0 \times 10^{19} \text{ cm}^{-2}$.

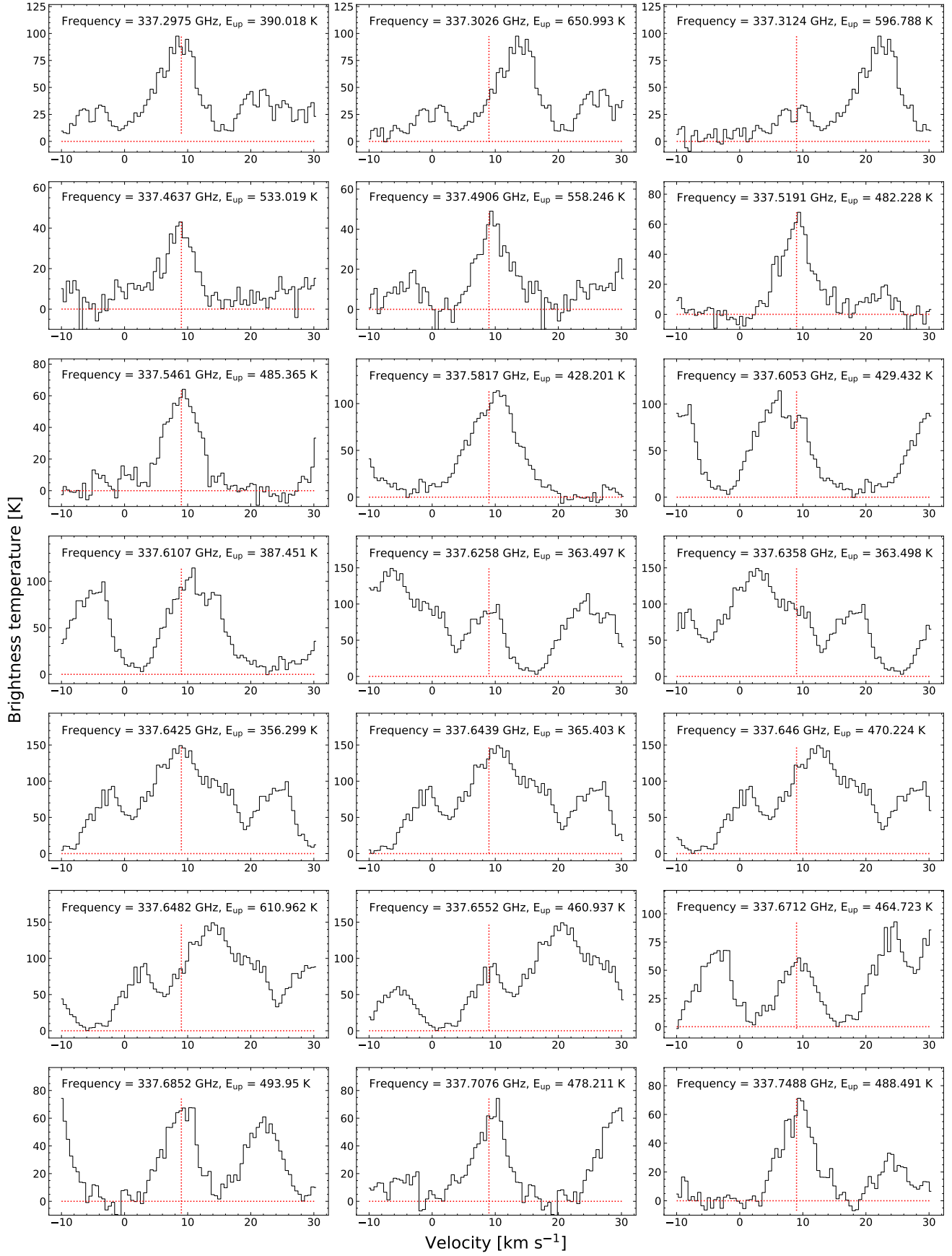


Fig. D.4. CH₃OH observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of ¹³CH₃OH with the ISM isotopologue ratio of ¹²C/¹³C = 77. The obtained column density was $1.0 \times 10^{19} \text{ cm}^{-2}$.

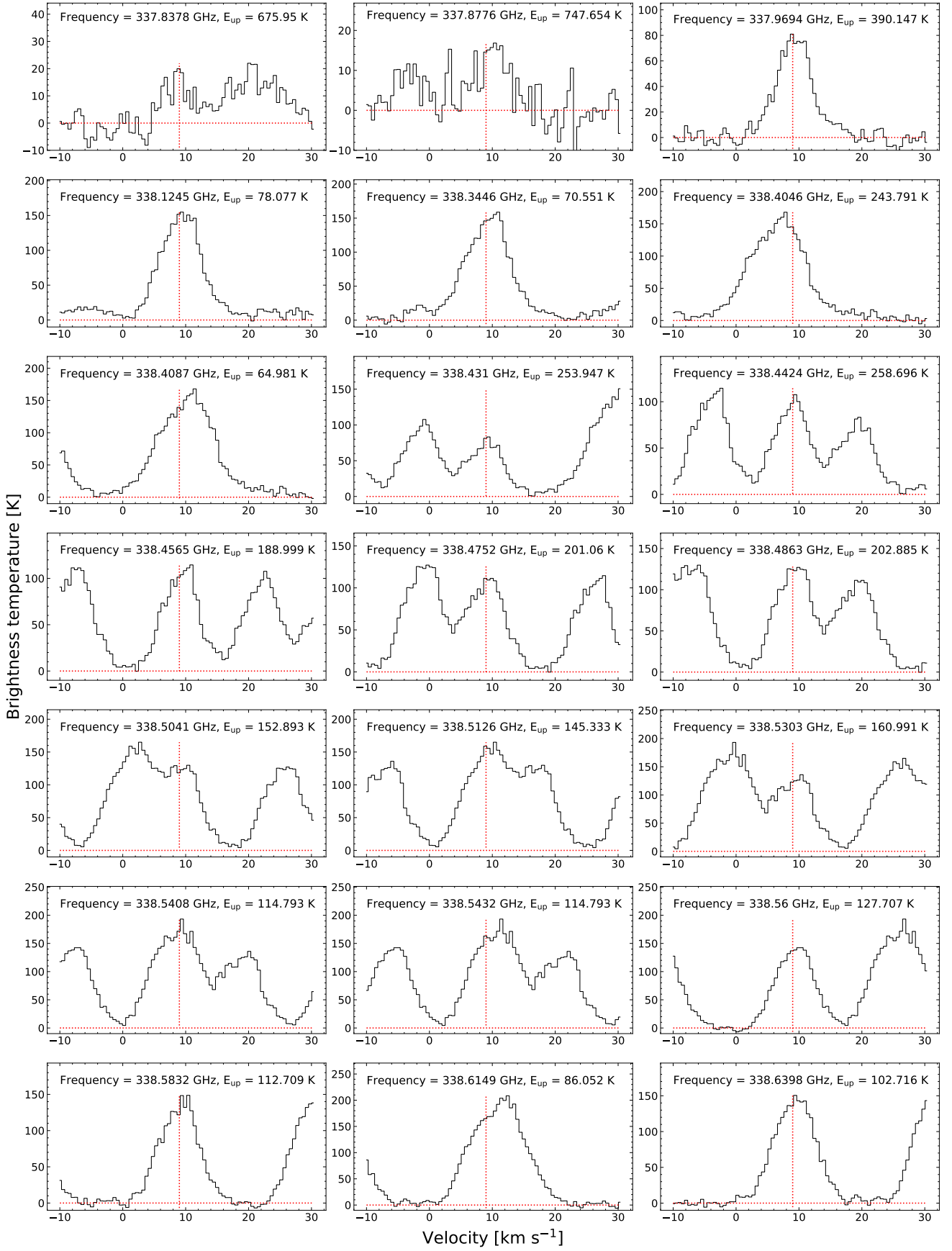


Fig. D.5. CH_3OH observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of $^{13}\text{CH}_3\text{OH}$ with the ISM isotopologue ratio of $^{12}\text{C}/^{13}\text{C} = 77$. The obtained column density was $1.0 \times 10^{19} \text{ cm}^{-2}$.

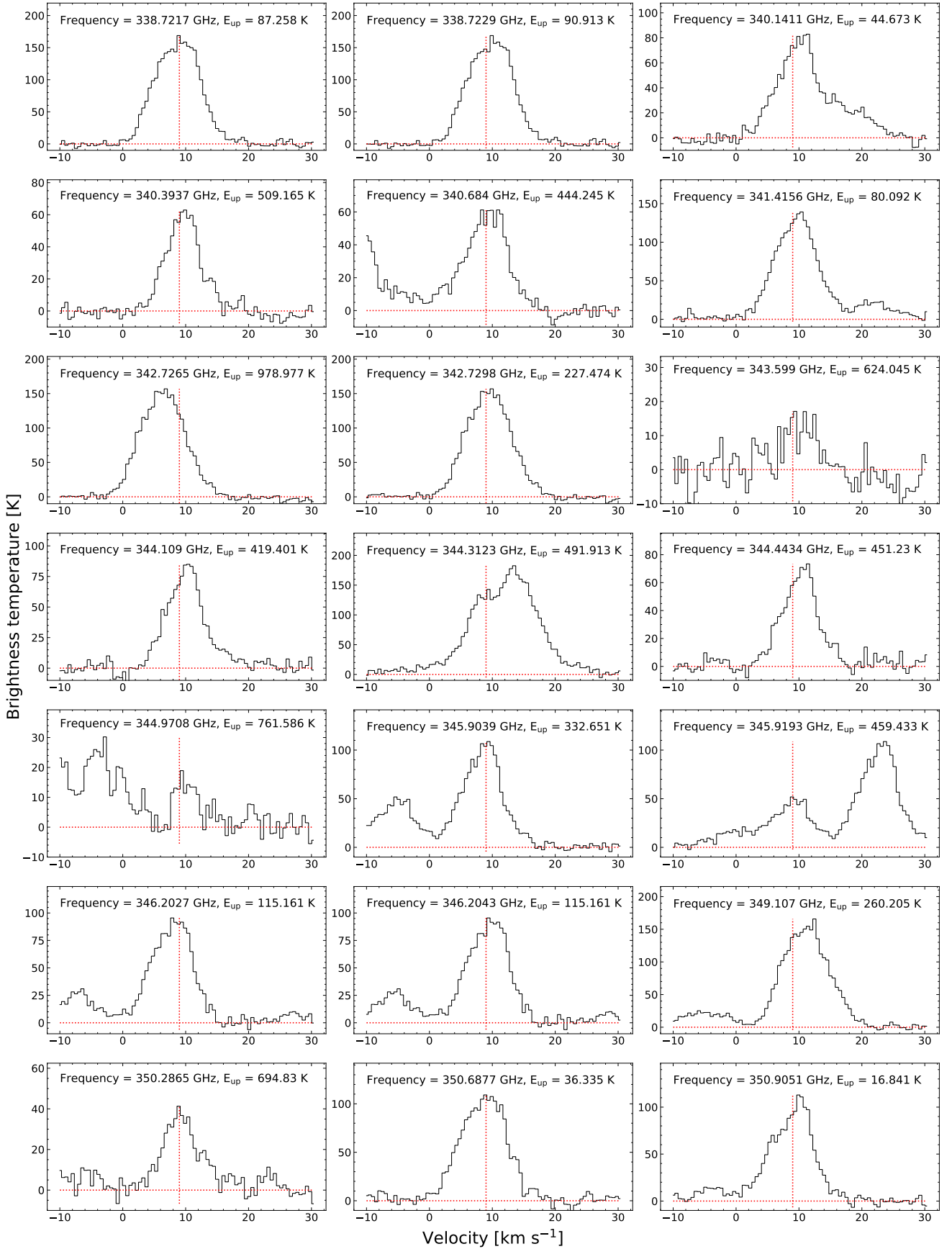


Fig. D.6. CH_3OH observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of $^{13}\text{CH}_3\text{OH}$ with the ISM isotopologue ratio of $^{12}\text{C}/^{13}\text{C} = 77$. The obtained column density was $1.0 \times 10^{19} \text{ cm}^{-2}$.

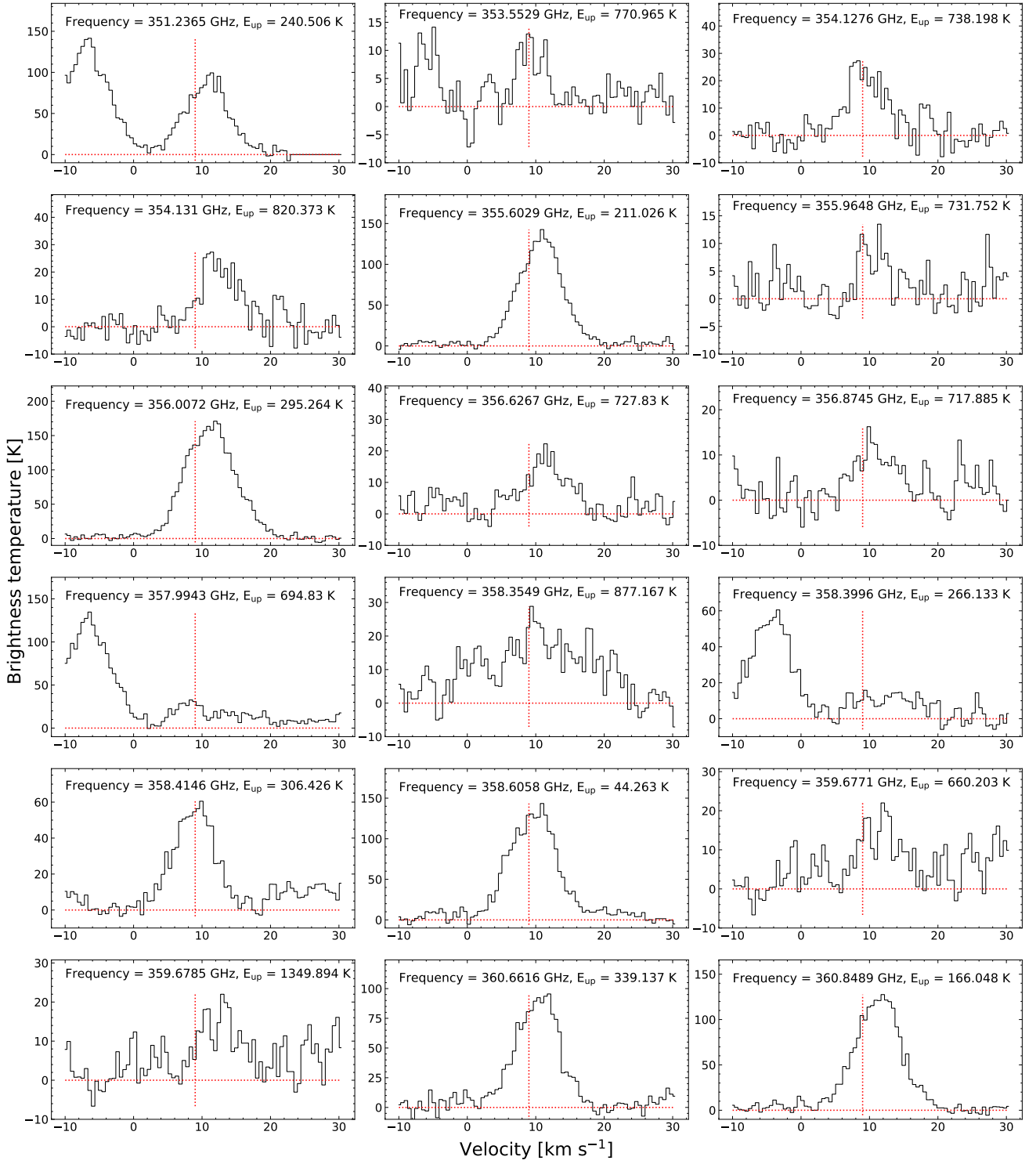


Fig. D.7. CH_3OH observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of $^{13}\text{CH}_3\text{OH}$ with the ISM isotopologue ratio of $^{12}\text{C}/^{13}\text{C} = 77$. The obtained column density was $1.0 \times 10^{19} \text{ cm}^{-2}$.

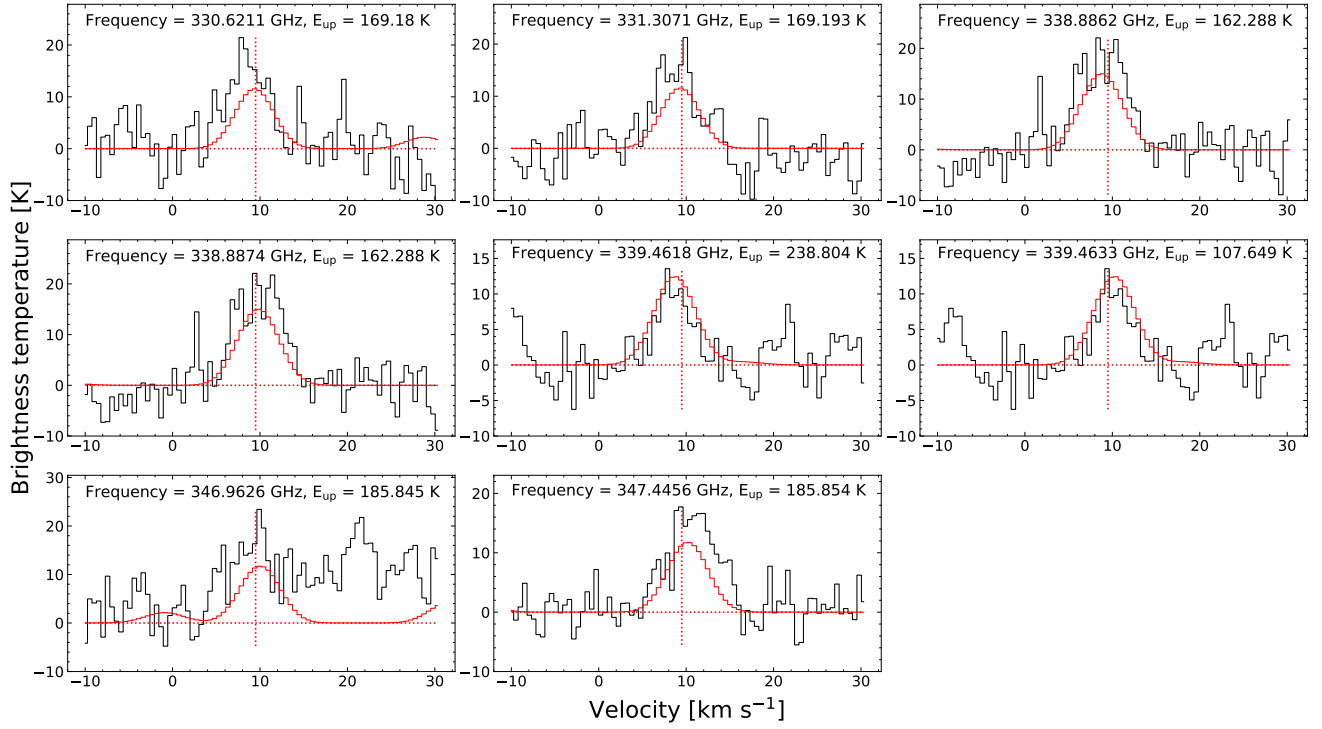


Fig. D.8. C₂H₅OH model lines and synthetic spectrum. The model fitted values obtained were column density = $6.7 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 130.0 K, line width (FWHM) = 5.0 km s⁻¹, and source velocity (v_{source}) = 9.5.

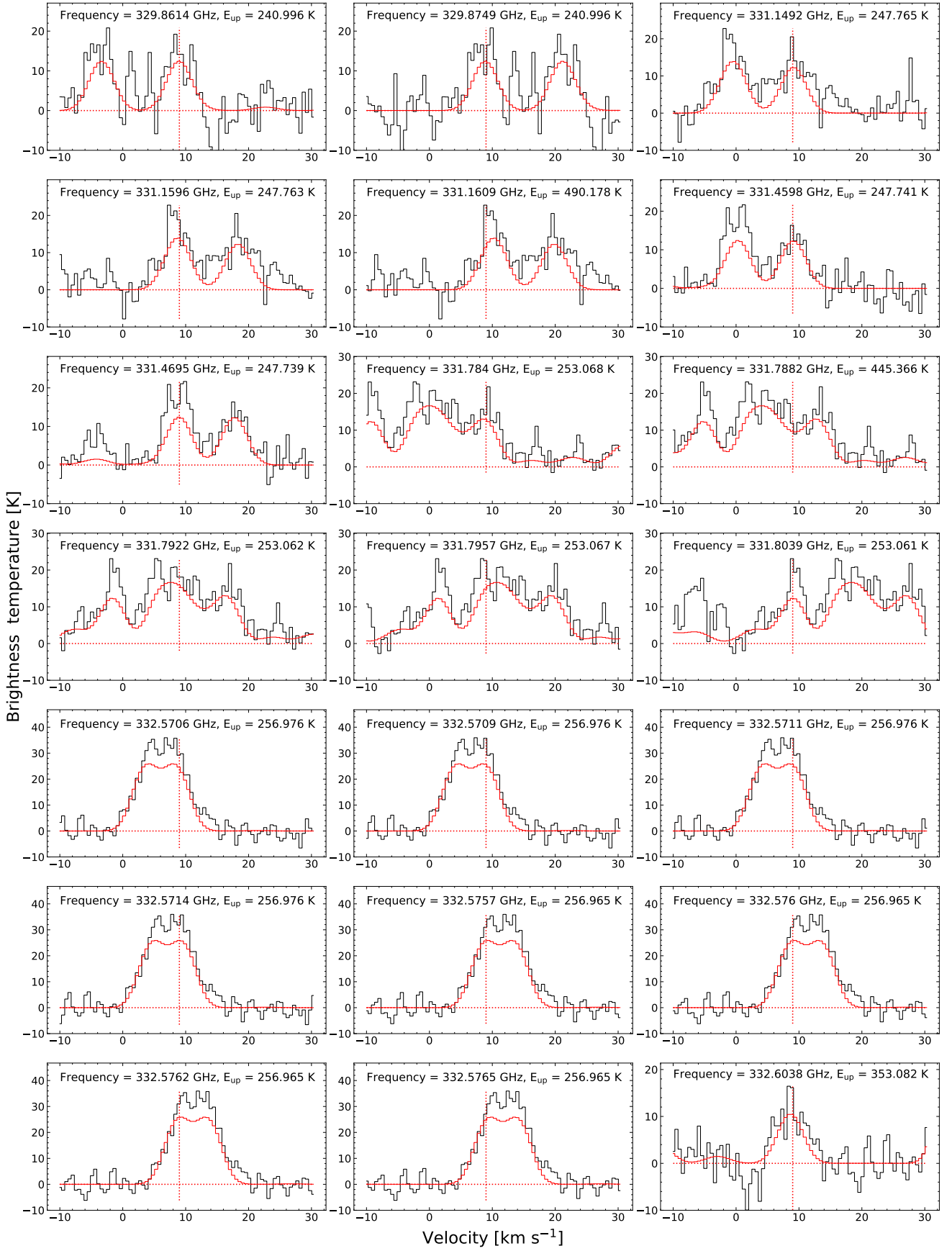


Fig. D.9. CH₃OCHO model lines and synthetic spectrum. The model fitted values obtained were column density = $1.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 4.5 km s⁻¹, and source velocity (v_{source}) = 9.0 km s⁻¹.

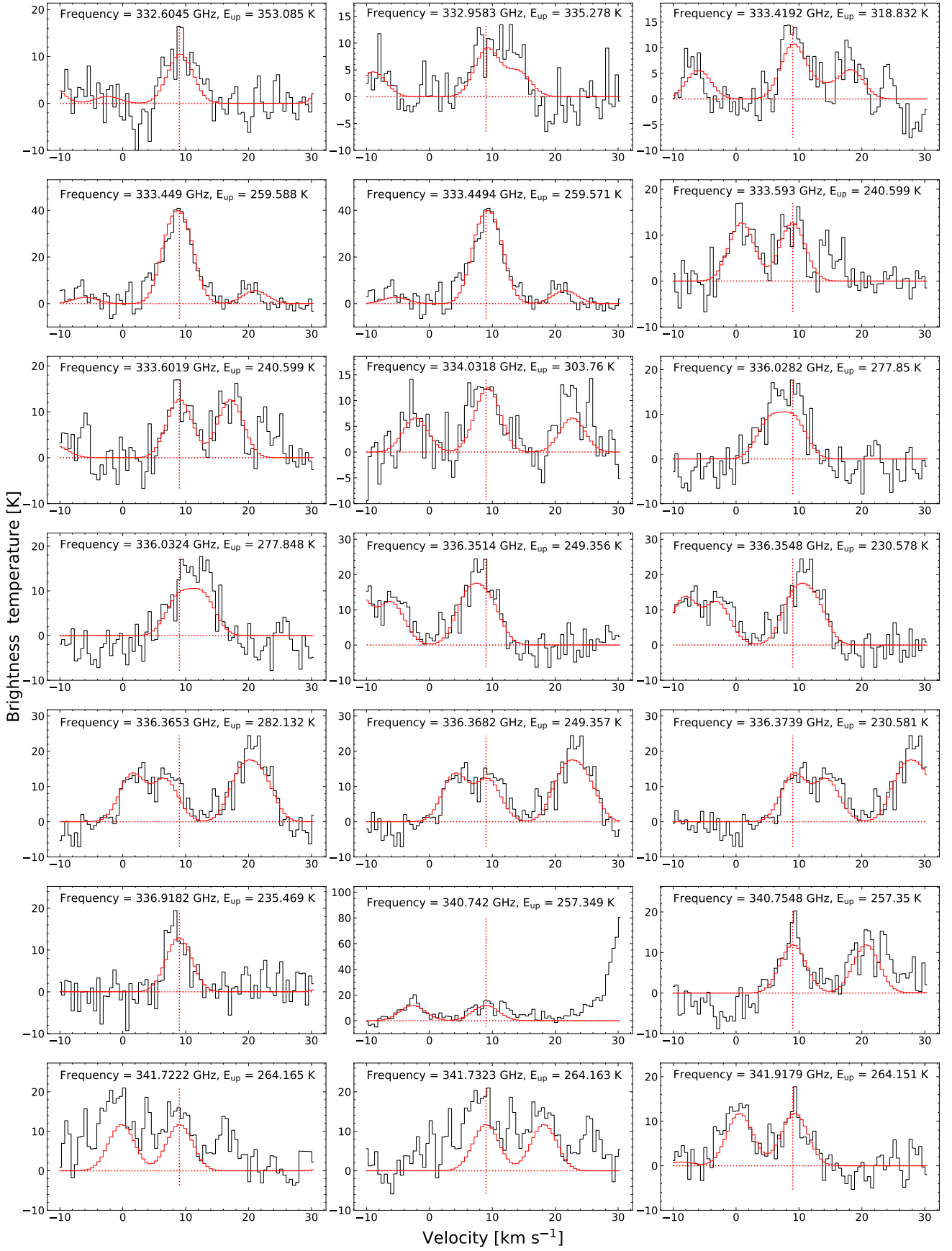


Fig. D.10. CH₃OCHO model lines and synthetic spectrum. The model fitted values obtained were column density = $1.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 4.5 km s⁻¹, and source velocity (v_{source}) = 9.0 km s⁻¹.

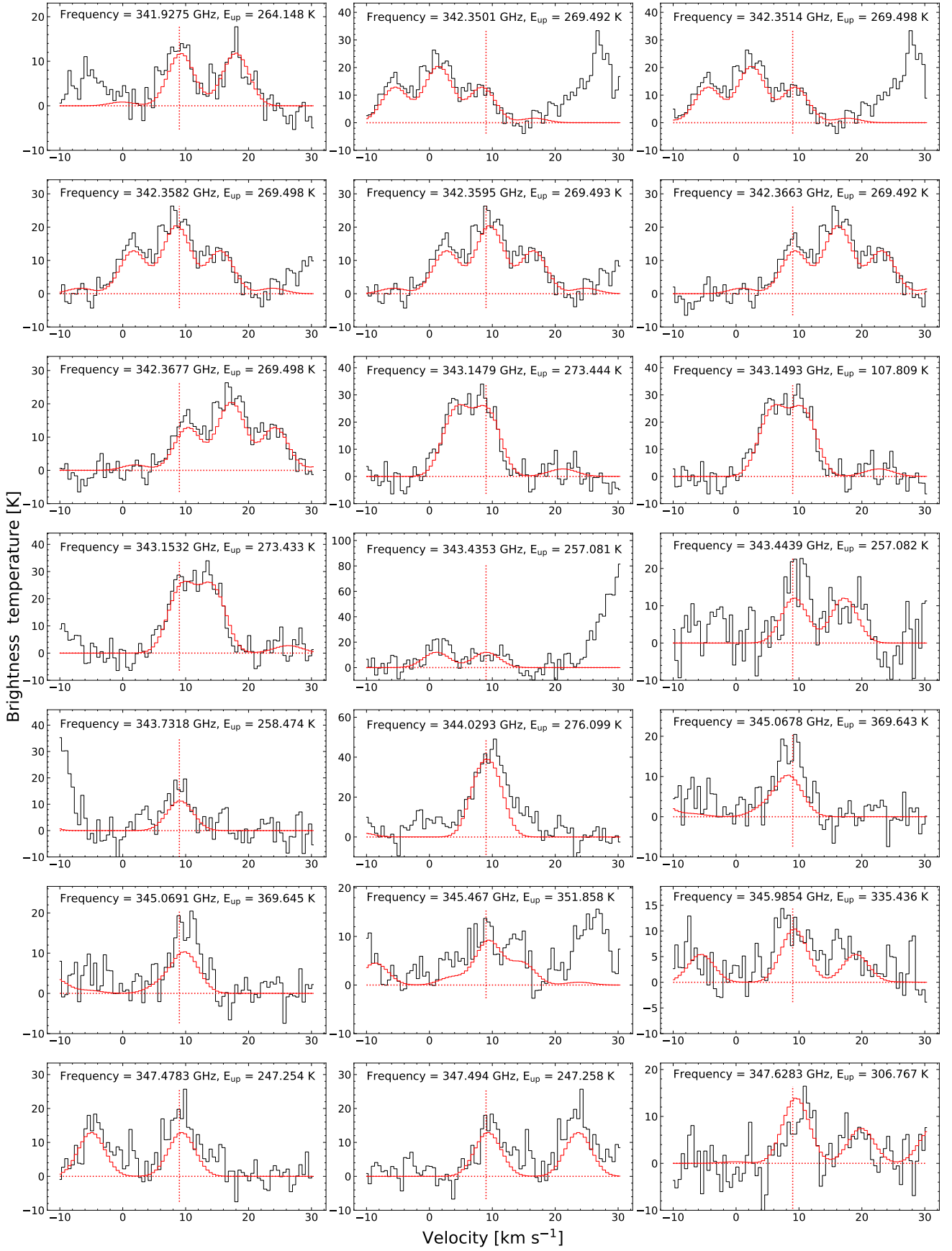


Fig. D.11. CH₃OCHO model lines and synthetic spectrum. The model fitted values obtained were column density = $1.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 4.5 km s⁻¹, and source velocity (v_{source}) = 9.0 km s⁻¹.

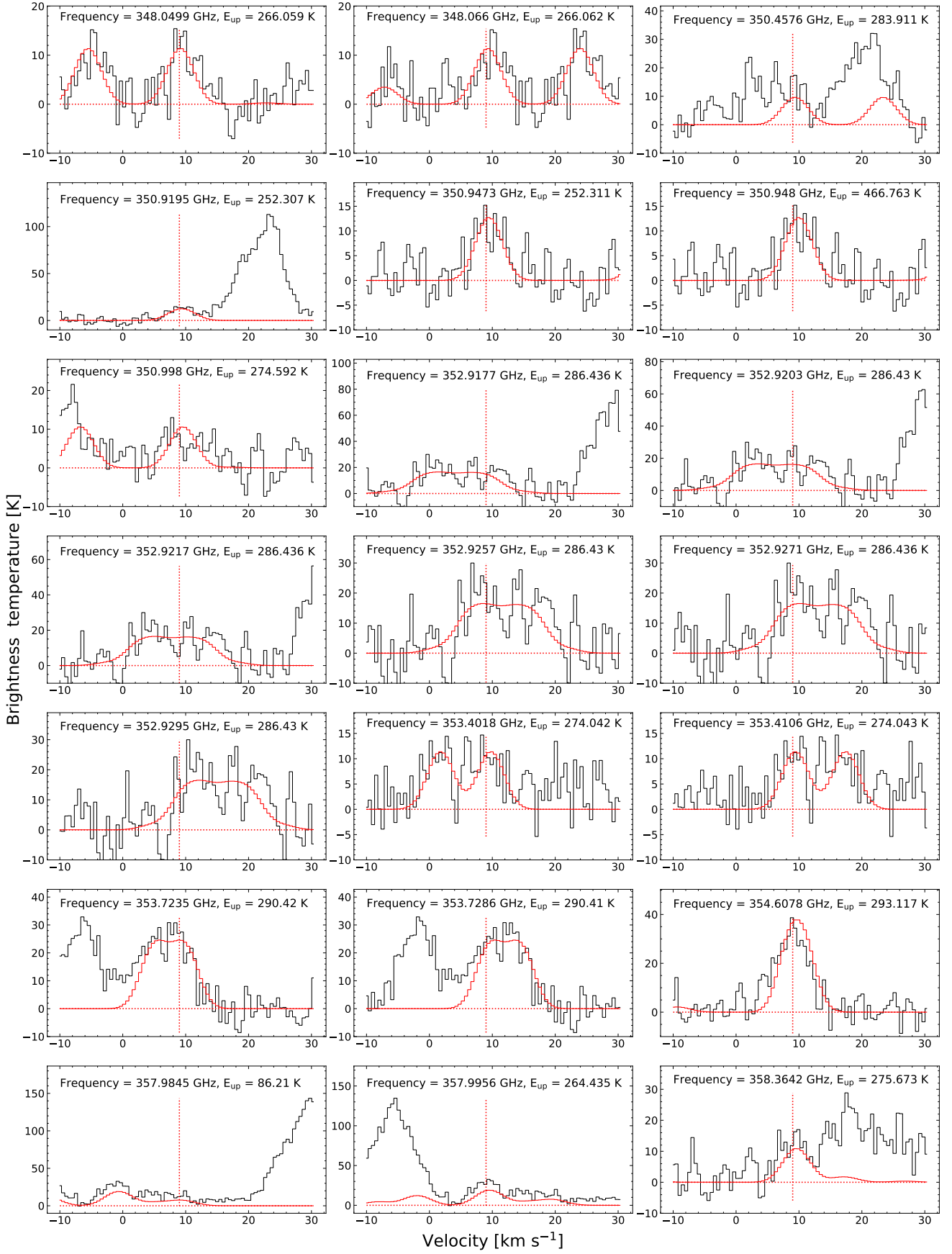


Fig. D.12. CH_3OCHO model lines and synthetic spectrum. The model fitted values obtained were column density = $1.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 4.5 km s^{-1} , and source velocity (v_{source}) = 9.0 km s^{-1} .

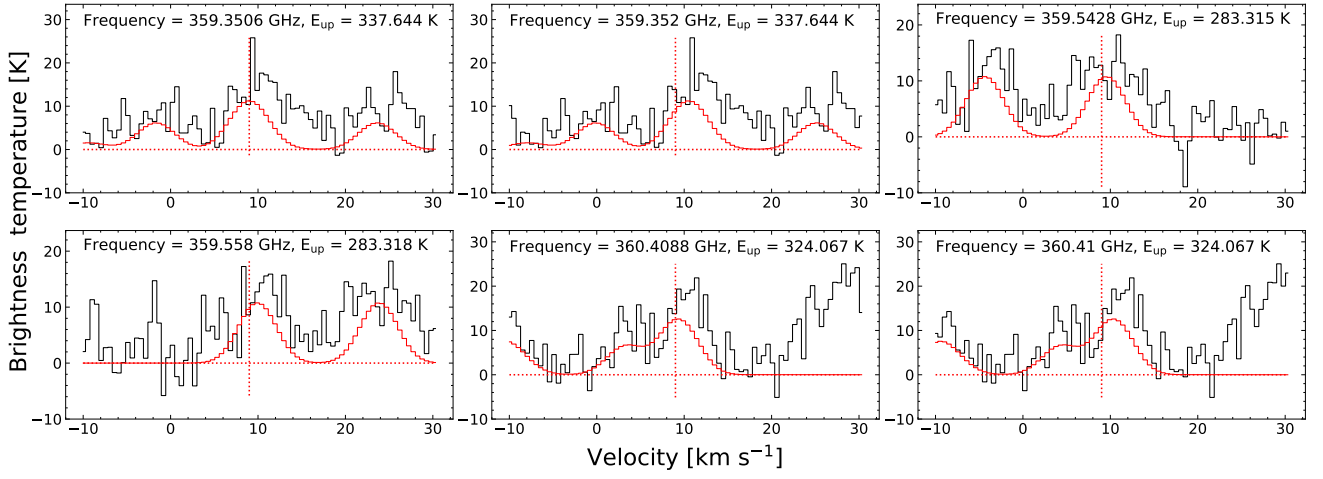


Fig. D.13. CH_3OCHO model lines and synthetic spectrum. The model fitted values obtained were column density = $1.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 4.5 km s^{-1} , and source velocity (v_{source}) = 9.0 km s^{-1} .

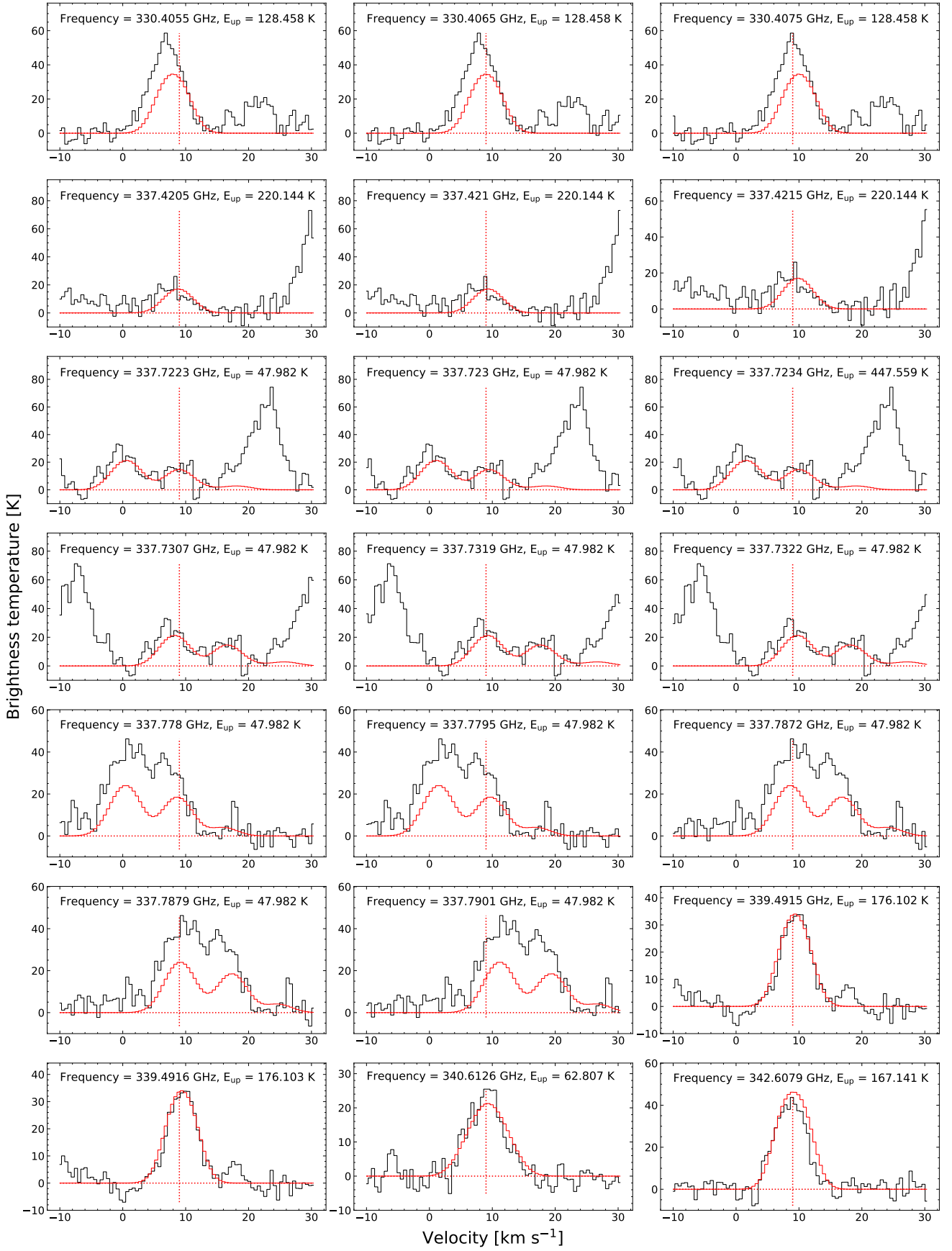


Fig. D.14. CH_3OCH_3 model lines and synthetic spectrum. The model fitted values obtained were column density = $2.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 5.0 km s⁻¹, and source velocity (v_{source}) = 9.0 km s⁻¹.

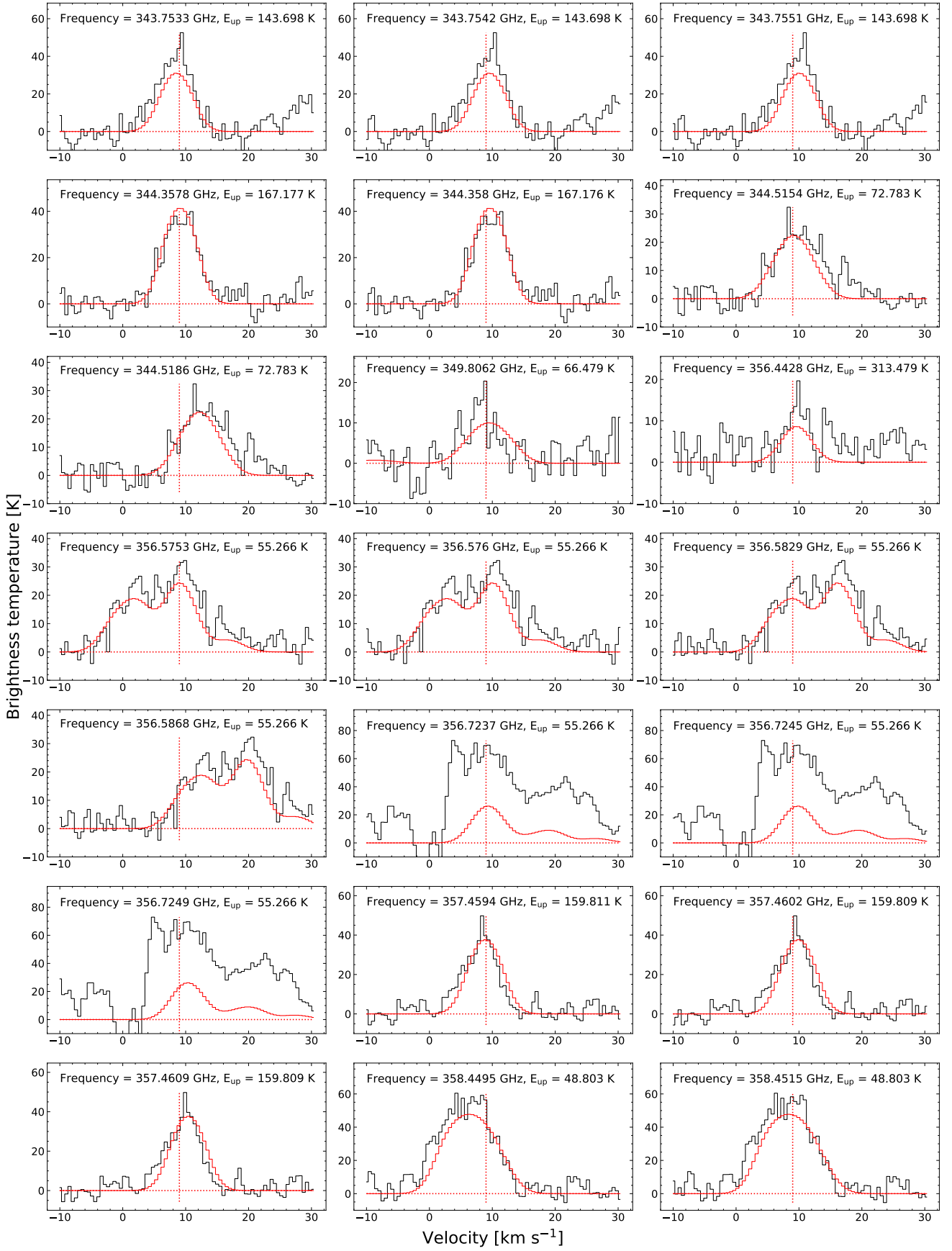


Fig. D.15. CH₃OCH₃ model lines and synthetic spectrum. The model fitted values obtained were column density = $2.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 5.0 km s⁻¹, and source velocity (v_{source}) = 9.0 km s⁻¹.

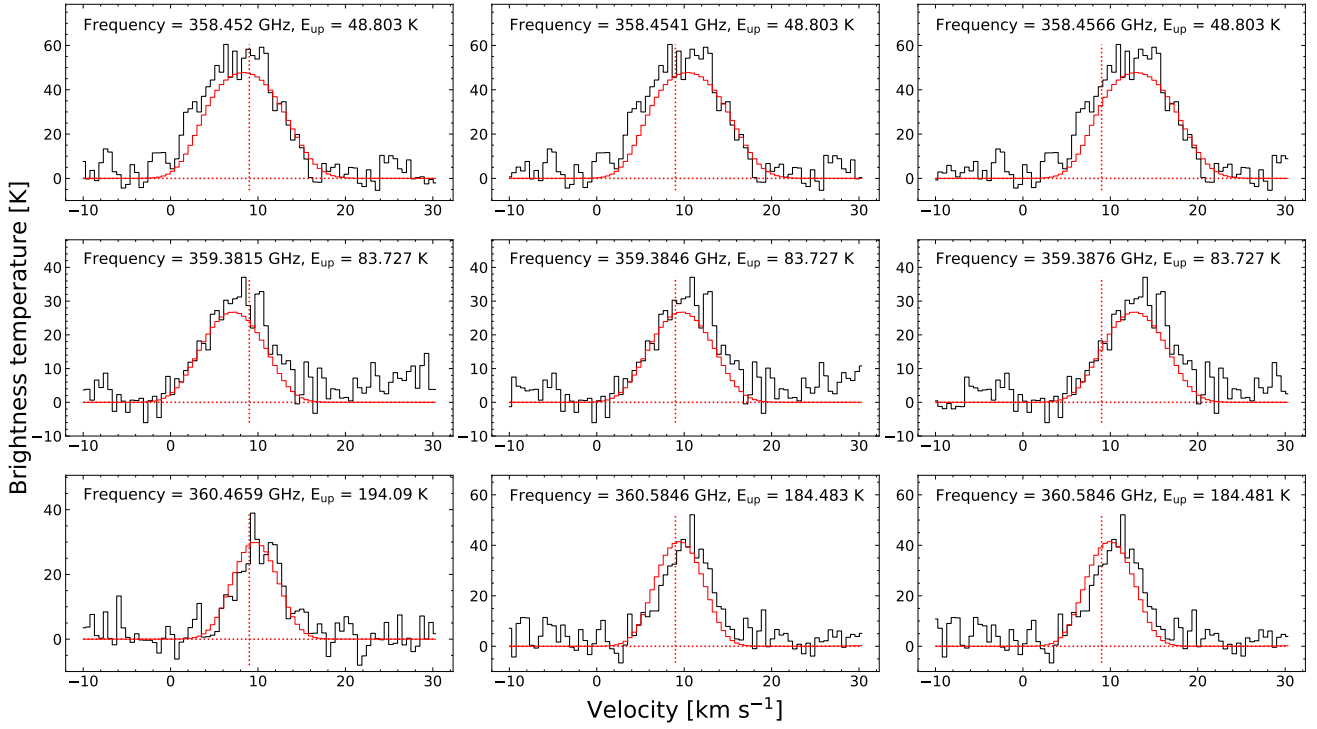


Fig. D.16. CH_3OCH_3 model lines and synthetic spectrum. The model fitted values obtained were column density = $2.4 \times 10^{17} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 110.0 K, line width (FWHM) = 5.0 km s^{-1} , and source velocity (v_{source}) = 9.0 km s^{-1} .

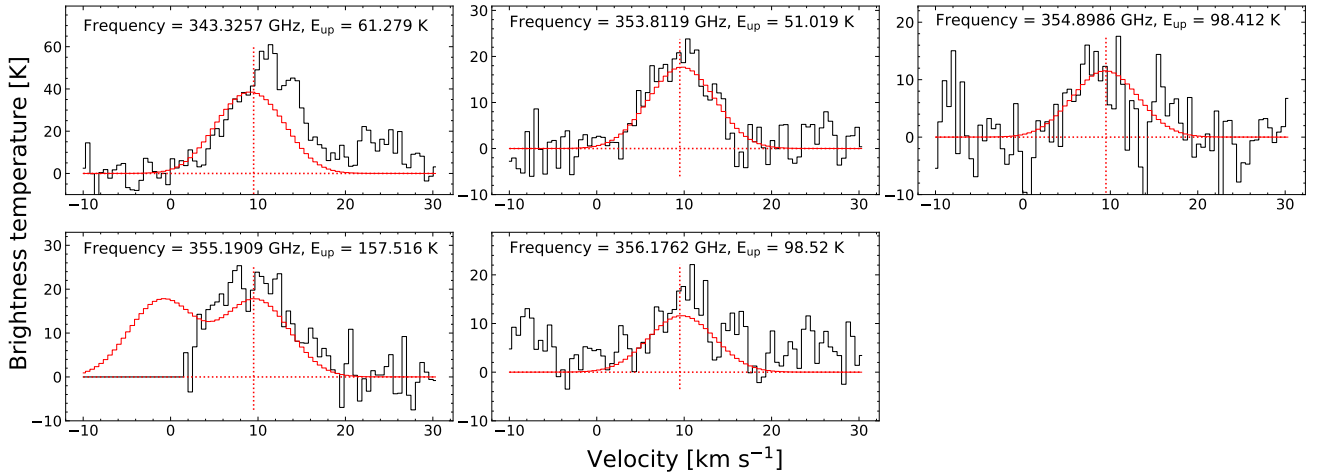


Fig. D.17. H_2^{13}CO model lines and synthetic spectrum. The model fitted values obtained were column density = $7.6 \times 10^{15} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 160.0 K, line width (FWHM) = 8.0 km s^{-1} , and source velocity (v_{source}) = 9.0 km s^{-1} .

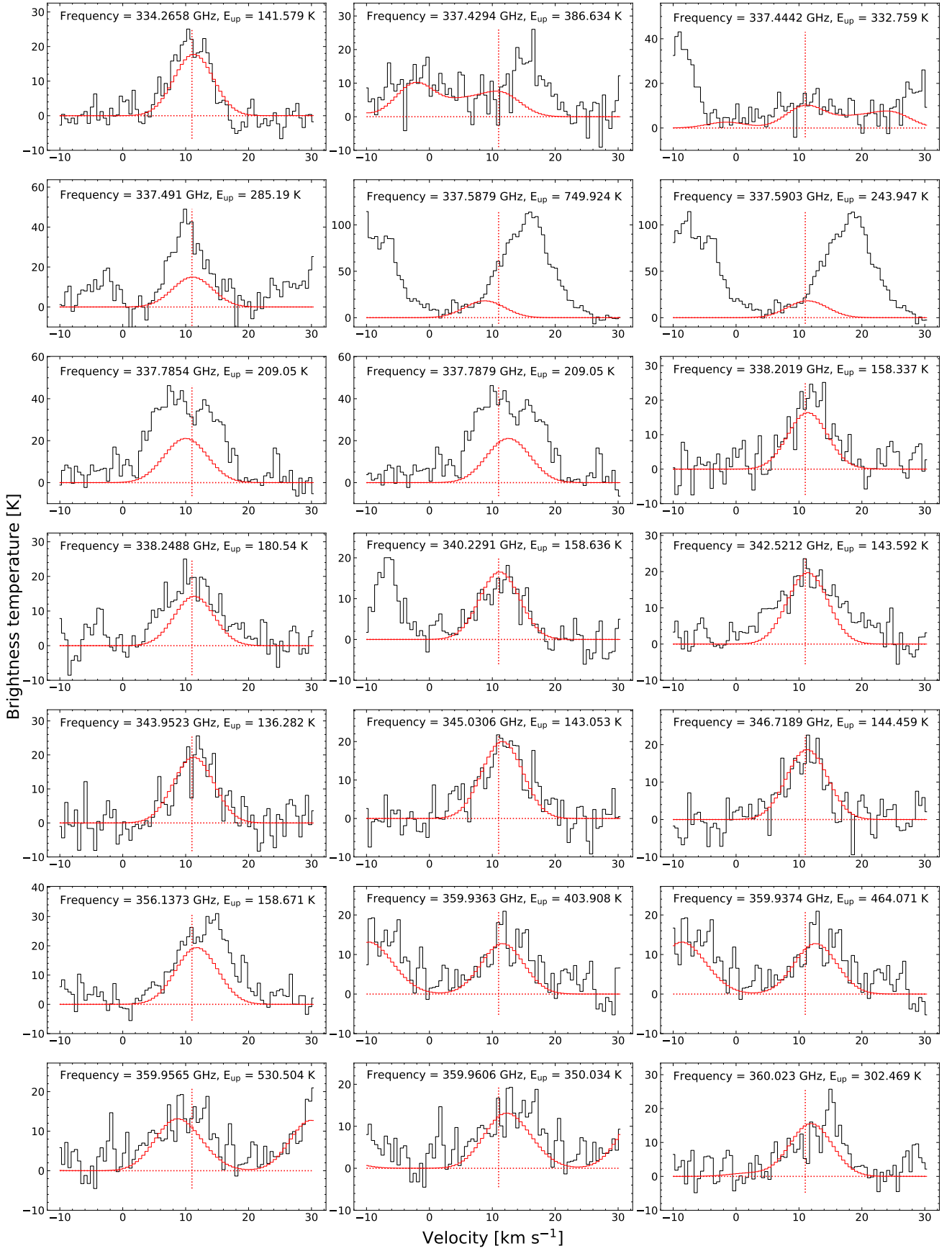


Fig. D.18. t-HCOOH model lines and synthetic spectrum. The model fitted values obtained were column density = $4.0 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 190.0 K, line width (FWHM) = 7.0 km s⁻¹, and source velocity (v_{source}) = 11.0 km s⁻¹.

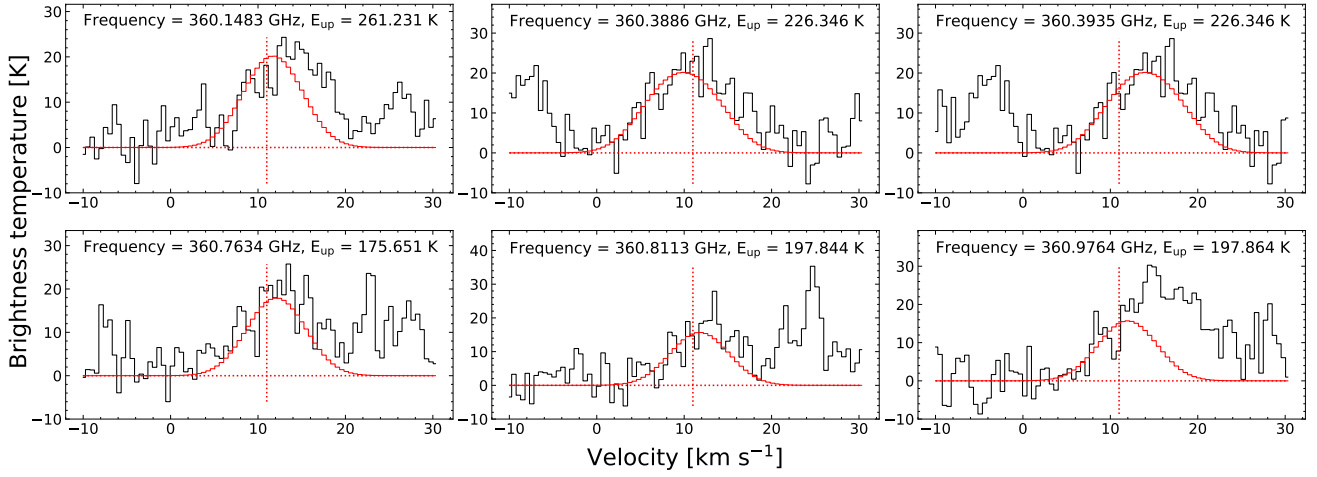


Fig. D.19. t-HCOOH model lines and synthetic spectrum. The model fitted values obtained were column density = $4.0 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 190.0 K, line width (FWHM) = 7.0 km s⁻¹, and source velocity (v_{source}) = 11.0 km s⁻¹.

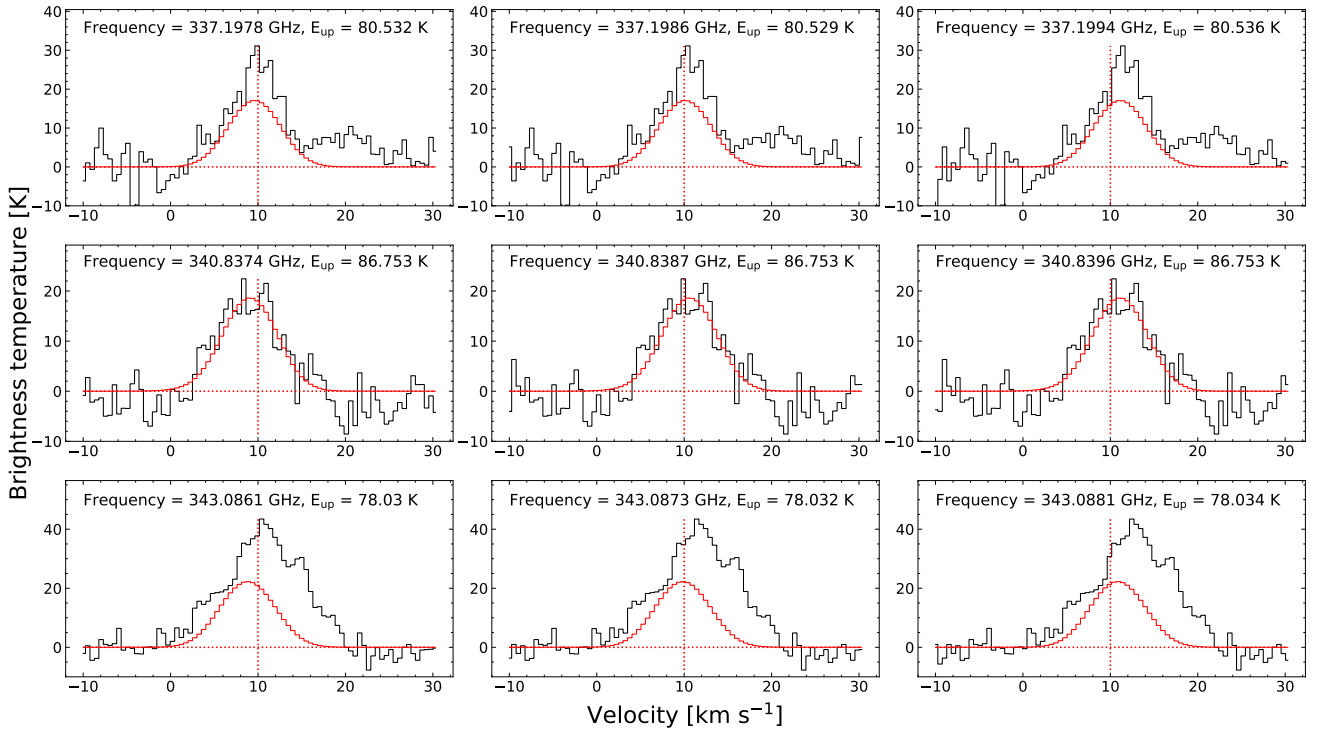


Fig. D.20. ³³SO model lines and synthetic spectrum. The model fitted values obtained were column density = $5.0 \times 10^{15} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 140.0 K, line width (FWHM) = 6.5 km s⁻¹, and source velocity (v_{source}) = 10.0 km s⁻¹.

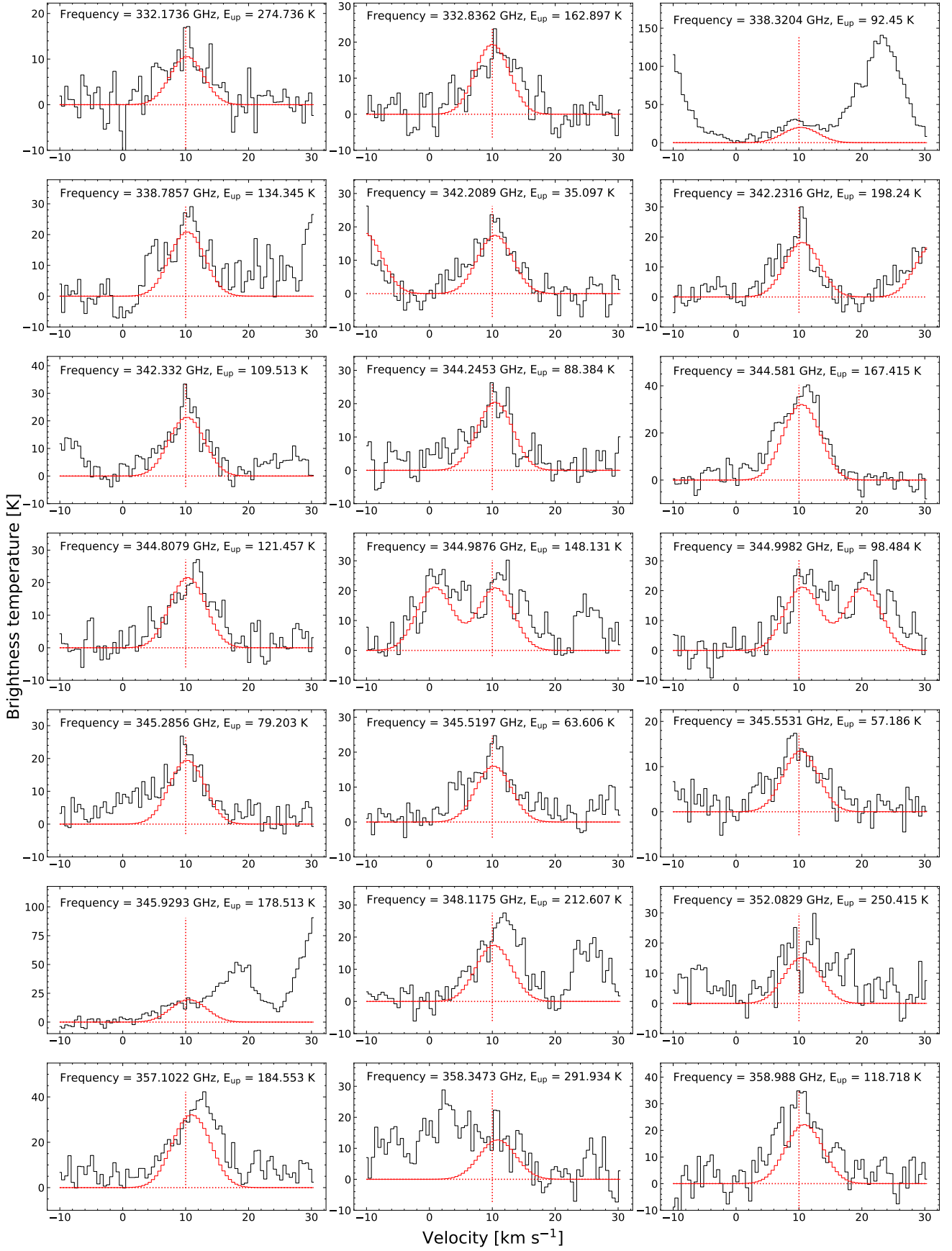


Fig. D.21. ³⁴SO₂ model lines and synthetic spectrum. The model fitted values obtained were column density = $3.3 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 130.0 K, line width (FWHM) = 6.0 km s⁻¹, and source velocity (v_{source}) = 10.0 km s⁻¹.

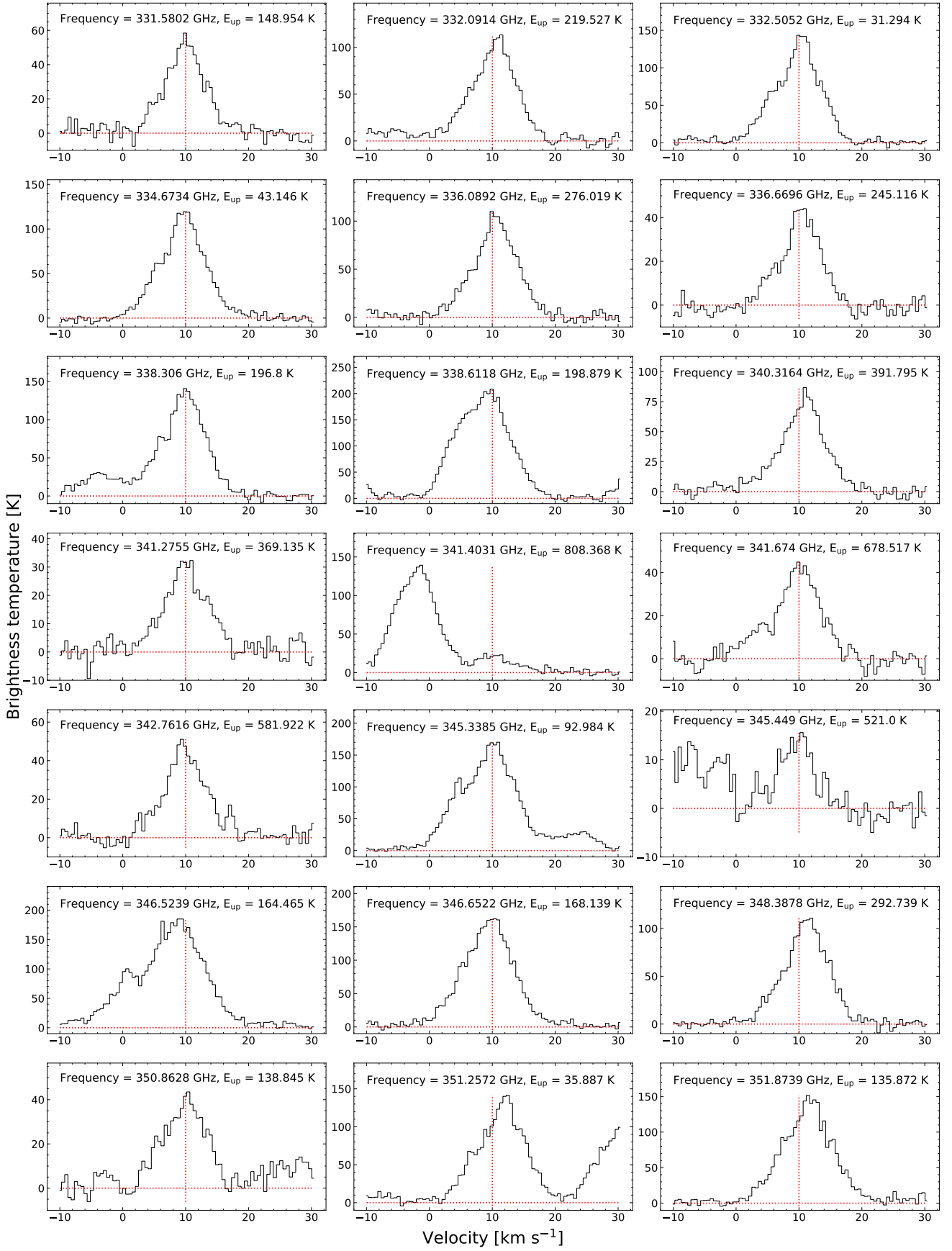


Fig. D.22. SO_2 observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of $^{34}\text{SO}_2$ with the ISM isotopologue ratio of $^{34}\text{S}/^{32}\text{S} = 22$. The obtained column density was $7.3 \times 10^{17} \text{ cm}^{-2}$.

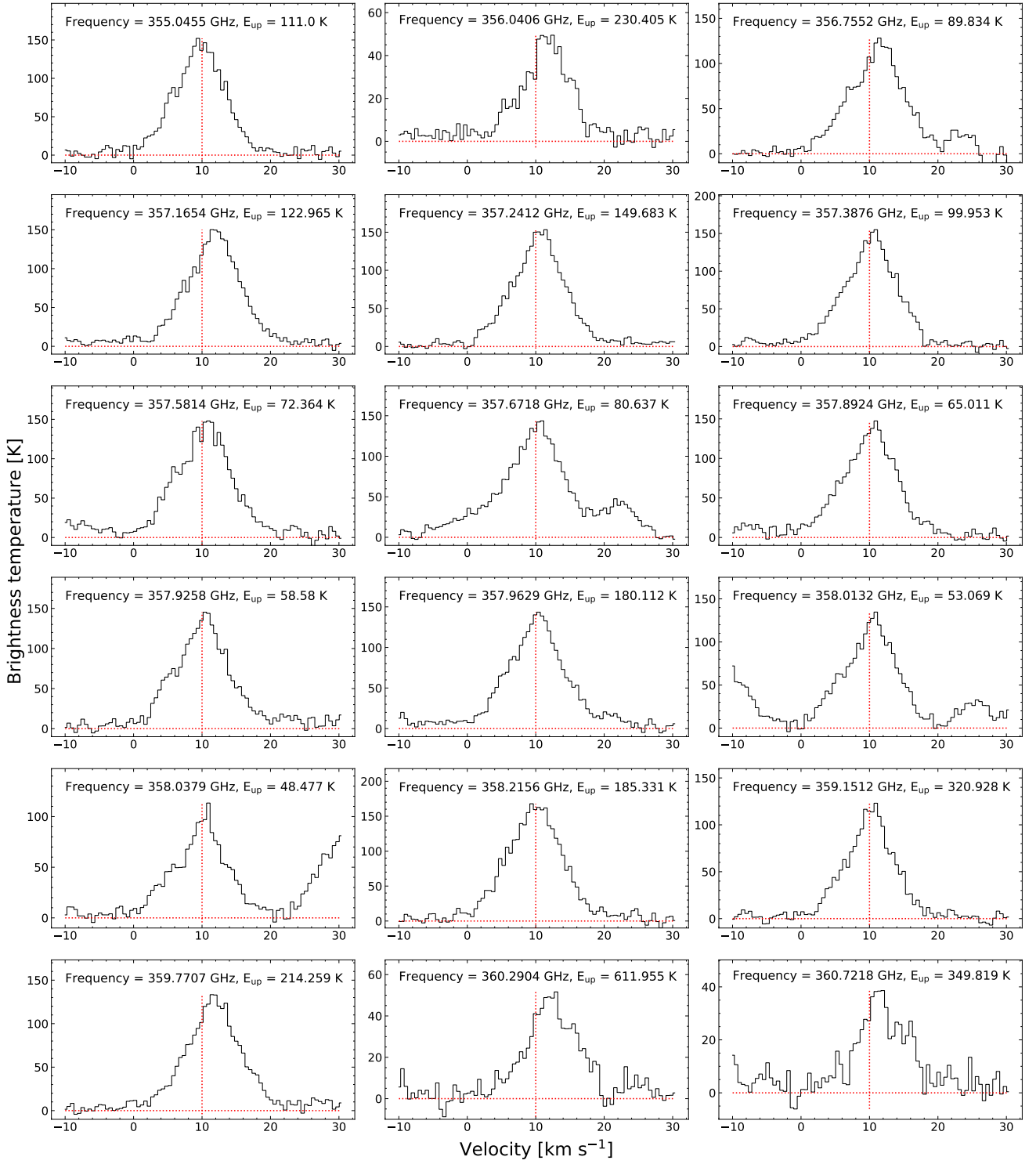


Fig. D.23. SO₂ observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of ³⁴SO₂ with the ISM isotopologue ratio of ³⁴S/³²S = 22. The obtained column density was $7.3 \times 10^{17} \text{ cm}^{-2}$.

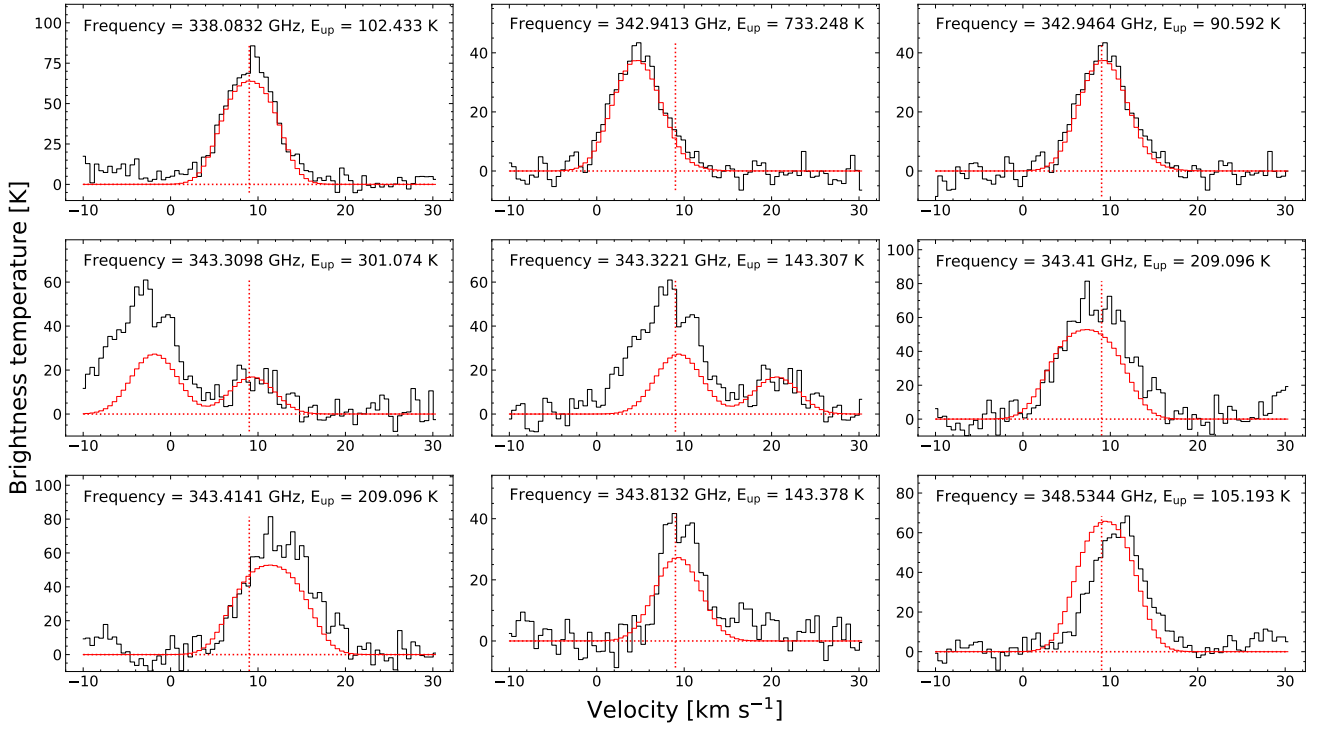


Fig. D.24. H₂CS model lines and synthetic spectrum. The model fitted values obtained were column density = $3.3 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 140.0 K, line width (FWHM) = 5.5 km s⁻¹, and source velocity (v_{source}) = 9.0 km s⁻¹.

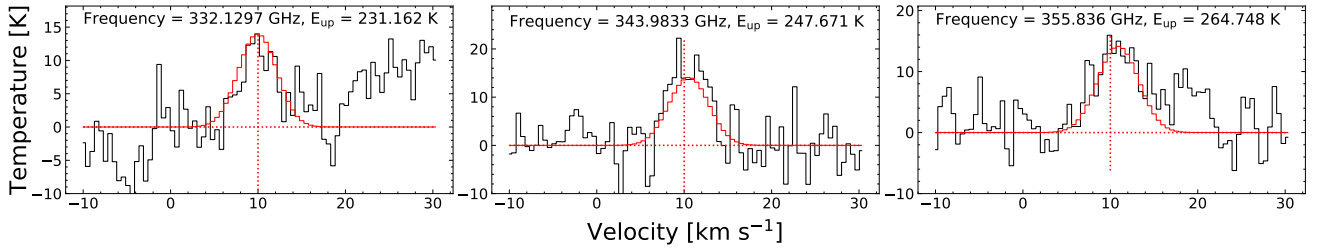


Fig. D.25. OC³⁴S model lines and synthetic spectrum. The model fitted values obtained were column density = $1.1 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 170.0 K, line width (FWHM) = 5.0 km s⁻¹, and source velocity (v_{source}) = 10.0 km s⁻¹.

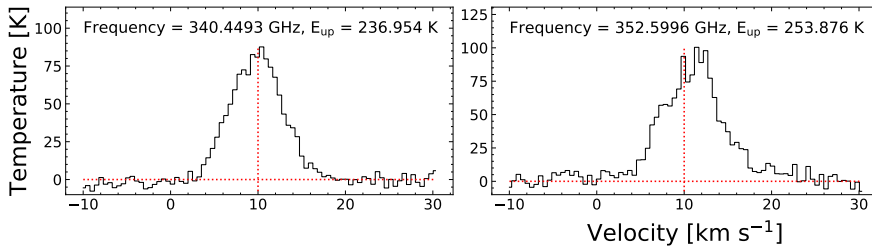


Fig. D.26. OCS observed spectral lines. In this case the column density was obtained by multiplying the fitted column density of OC³⁴S with the ISM isotopologue ratio of $^{34}\text{S}/^{32}\text{S} = 22$. The obtained column density was $2.4 \times 10^{17} \text{ cm}^{-2}$.

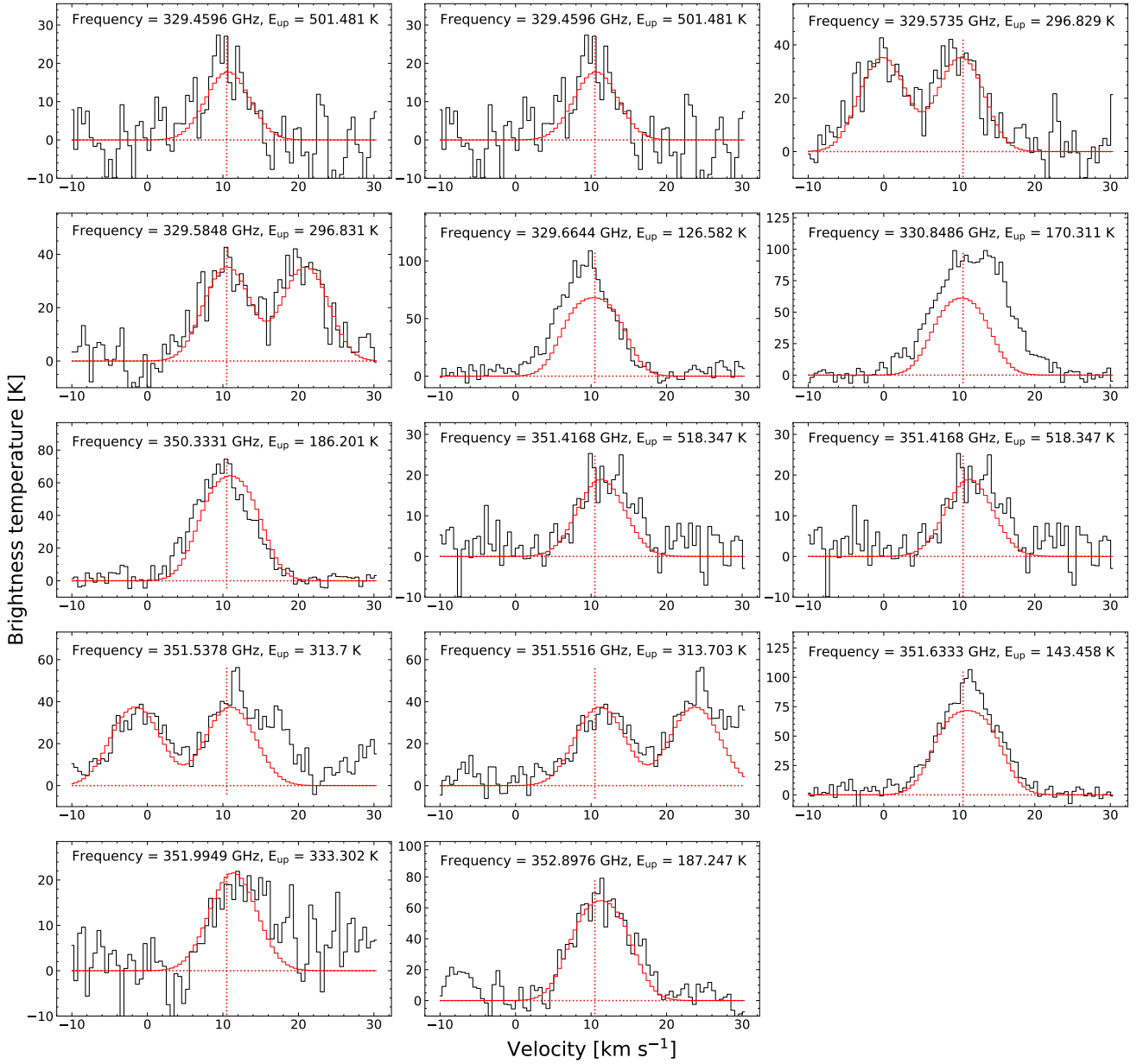


Fig. D.27. HNC model lines and synthetic spectrum. The model fitted values obtained were column density = $5.0 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 150.0 K, line width (FWHM) = 6.5 km s⁻¹, and source velocity (v_{source}) = 10.5 km s⁻¹.

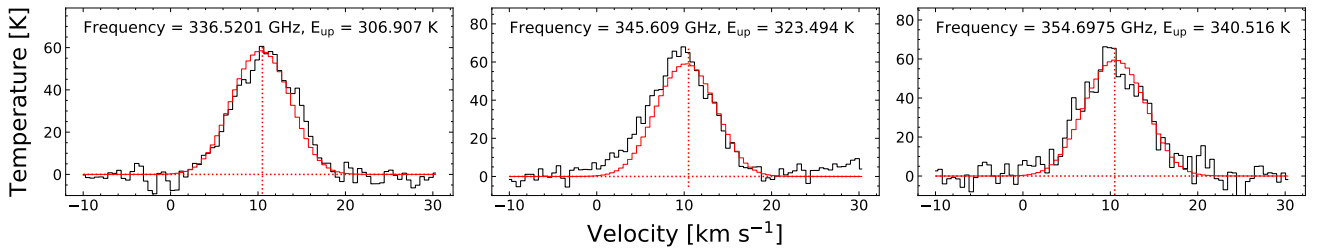


Fig. D.28. HC₃N model lines and synthetic spectrum. The model fitted values obtained were column density = $4.0 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 210.0 K, line width (FWHM) = 7.0 km s⁻¹, and source velocity (v_{source}) = 10.0 km s⁻¹.

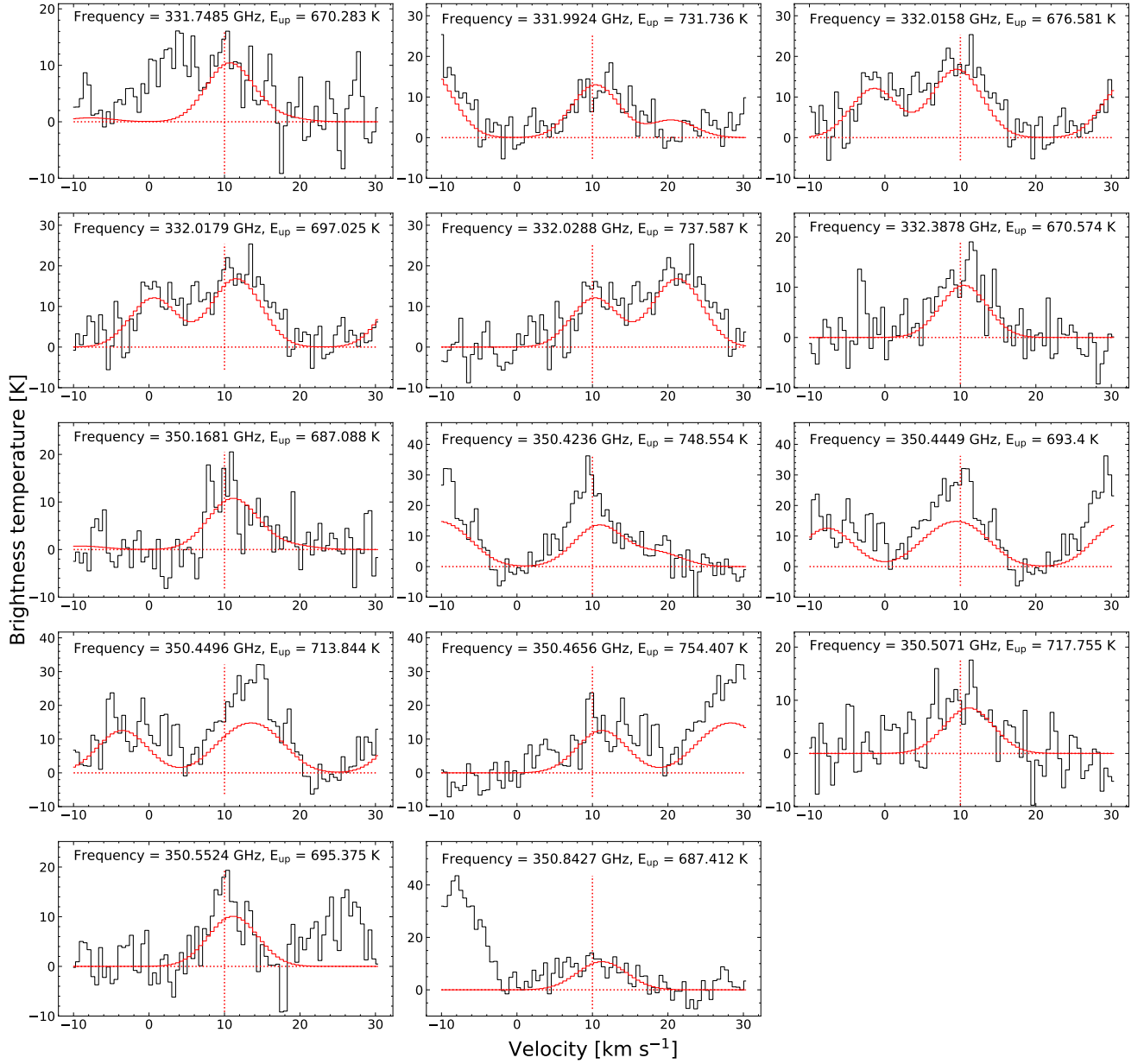


Fig. D.29. CH₃CN, v8=1 model lines and synthetic spectrum. The model fitted values obtained were column density = $5.5 \times 10^{16} \text{ cm}^{-2}$, excitation temperature (T_{ex}) = 140.0 K, line width (FWHM) = 7.0 km s⁻¹, and source velocity (v_{source}) = 10.5 km s⁻¹.

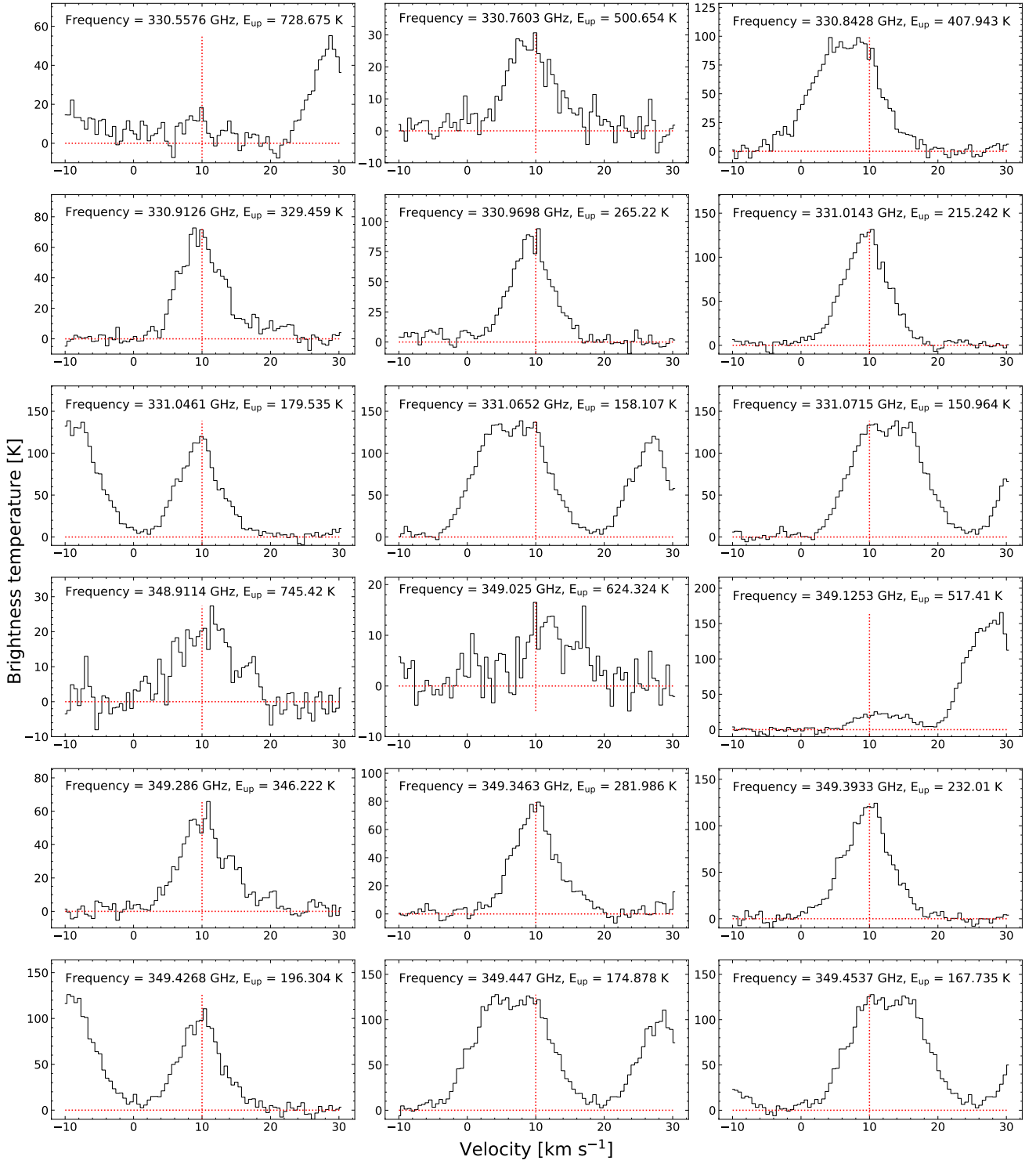


Fig. D.30. CH_3CN , $v=0$ observed spectral lines. In this case the column density was obtained using the higher energy $v = 1$ transition. The obtained column density was $5.5 \times 10^{16} \text{ cm}^{-2}$.